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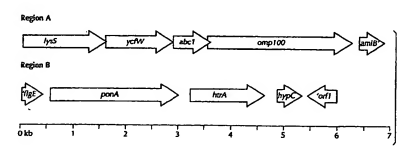
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(54) Lawsonia intracellularis proteins, and related methods and materials

(57) Isolated polynucleotide molecules contain a nucleotide sequence that encodes a *L. intracellularis* HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein, a substantial portion of the sequences, or a homol-

ogous sequence. Related polypeptides, immunogenic compositions and assays are described.

FIG. 1



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Description

FIELD OF THE INVENTION

[0001] The present invention relates to protens derived from Lawsonia intracellularis and encompasses related proteins, nucleic acids, and immunogenic compositions. The immunogenic compositions are particularly useful in prevention of L. intracellularis infections in susceptible animals, such as pigs. The proteins, fragments, and nucleic acids can also be employed as diagnostic agents.

10 BACKGROUND OF THE INVENTION

[0002] Commercially raised pigs are sensitive to a wide spectrum of intestinal diseases or syndromes that are collectively referred to as porcine proliferative enteropathy (PPE). These diseases include intestinal adenomatosis complex (Barker I. K. et al., 1985, In "Pathology of Domestic Animals," 3rd Edition, Vol. 2 p. 1-237, eds. K. V. F. Jubb et al. (Academic Press: Orlando)), porcine intestinal adenomatosis (PIA), necrotic enteritis (Rowland A. C. et al., 1976, *Veterinary Record* 97:-178-180), proliferative haemorrhagic enteropathy (Love, R. J. et al., 1977, *Veterinary Record* 100: 473), regional ileitis (Jonsson, L. et al., 1976, Acta *Veterinaria Scandinavica* 17: 223-232), haemorrhagic bowel syndrome (O'Neil, I. P. A., 1970, *Veterinary Record* 87:742-747), porcine proliferative enteritis and *Campylobacter* sppinduced enteritis (Straw, B. E., 1990, *Journal of American Veterinary Medical Association* 197: 355-357).

[0003] One major type of PPE is non-haemorrhagic and is manifested by porcine intestinal adenomatosis (PIA). This form of PPE frequently causes growth retardation and mild diarrhea. Another important type of PPE is haemorrhagic. It is often fatal, and is manifested by proliferative haemorrhagic enteropathy (PHE) wherein the distal small intestine lumen becomes engorged with blood.

[0004] While PPE in pigs is commercially most important, PPE is also a problem in the raising of hamsters (Stills, H. F., 1991, *Infection and Immunology* 59: 3227-3236), ferrets (Fox et al., 1989, *Veterinary Pathology* 26: 515-517), guinea pigs (Elwell et al., 1981, Veterinary Pathology 18: 136-139), rabbits (Schodeb et al., 1990, *Veterinary Pathology* 27: 73-80) and certain birds (Mason *et al.*, 1998).

[0005] The organism that causes PPE is the Campylobacter-like bacterium "L. intracellularis" (McOrist S et al, 1995, International Journal Of Systematic Bacteriology 45: 820-825). This organism is also known as heal symbiont intracellularis (Stills, 1991, supra). PPE-hike diseases in pigs may also be caused by other species of Campylobacter (Gebhart et al., 1983, American Journal of Veterinary Research 44: 361-367).

[0006] L. intracellularis is located in the cytoplasm of villi and intestinal crypt cells of infected animals, where it causes structural irregularities and enterocyte proliferation. Abscesses form as the villi and intestinal crypts become branched and fill with inflammatory cells.

[0007] Current control of PPE relies on the use of antibacterial compounds. There is, however, a need for alternative means of controlling *L. intracellularis* infection.

[0008] International Patent Application No. PCT/AU96/00767 describes *L. intracellularis* polypeptides and immunogenic compositions that are useful as vaccines. There is, however, a need for additional compositions that confer resistance to *L. intracellularis* infection.

SUMMARY OF THE INVENTION

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[0009] The present invention relates to an isolated polynucleotide molecule comprising a nucleotide sequence that is selected from the group consisting of:

- a) a nucleotide sequence encoding L. intracellularis HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein;
- b) a nucleotide sequence that is a substantial part of the nucleotide sequence encoding the *L. intracellularis* HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein; and
- c) a nucleotide sequence that is homologous to the nucleotide sequence of a) or b).

[0010] In another aspect, the invention relates to a recombinant vector comprising these polynucleotide molecules, including those encoding a carrier or fusion partner such that expression of the recombinant vector results in a fusion protein comprising the carrier or fusion partner fused to a protein or polypeptide encoded by the nucleotide sequences described above. The invention also encompasses transformed host cells comprising these recombinant vectors and polypeptides produced by such transformed host cells.

[0011] In another aspect, the present invention relates to an isolated polypeptide that is selected from the group consisting of:

- (a) L. intracellularis HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein;
- (b) a polypeptide having an amino acid sequence that is homologous to that of the *L. intracellularis* HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein;
- (c) a polypeptide consisting of a substantial portion of the *L. intracellularis* HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein or of the polypeptide having an amino acid sequence that is homologous to that of the *L. intracellularis* HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein;
- (d) a fusion protein comprising the protein or polypeptide of (a), (b) or (c) fused to another protein or polypeptide; and
- (e) an analog or derivative of the protein or polypeptide of (a), (b), (c) or (d).

[0012] The present invention further provides a polynucleotide molecule comprising a nucleotide sequence of greater than 20 nucleotides having promoter activity and found within SEQ ID NO: 2 from about nt 2691 to about nt 2890.

[0013] The present invention further relates to a method of preparing any of these polypeptides, comprising culturing host cells transformed with a recombinant expression vector and recovering the expressed polypeptide from the cell culture. The vector comprises a polynucleotide molecule comprising a nucleotide sequence encoding any of the polypeptides, the nucleotide sequence being in operative association with one or more regulatory elements. Culturing is conducted under conditions conducive to expression of the polypeptide.

[0014] In yet another aspect, the invention relates to an isolated antibody that specifically reacts with any of the *L. intracellularis* HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 proteins or polypeptides described above.

[0015] The invention also relates to an immunizing composition that comprises an immunologically effective amount of a protein, polypeptide, antibody, or polynucleotide of the invention in combination with a pharmaceutically acceptable carrier. The present invention encompasses a method of immunizing a PPE susceptible animal against *L. intracellularis* infection that comprises administering to the animal the immunizing composition.

[0016] The invention also relates to a kit for immunizing a PPE susceptible animal against a disease condition caused or exacerbated by *L. intracellularis* that comprises a container having therein an immunologically effective amount of one of the proteins, polypeptides, antibodies, or polynucleotides described above. The invention also relates to a kit for detecting the presence of *L. intracellularis*, an *L. intracellularis* specific amino acid or nucleotide sequence, or an anti- *L. intracellularis* antibody, comprising a container that has therein a protein, polypeptide, polynucleotide, or antibody of the invention.

BRIEF DESCRIPTION OF DRAWINGS

[0017]

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Figure 1 shows the arrangement of gene cluster A, containing the genes encoding the LysS, YcfW, ABC1 and Omp100 proteins, and the arrangement of gene cluster B, encoding the PonA, HtrA, and HypC proteins.

Figure 2 shows an alignment of the YcfW amino acid sequence with the most similar sequence found in a search of the GenBank database.

Figure 3 shows an alignment of the ABC1 amino acid sequence with the most similar sequence found in a search of the GenBank database.

Figure 4 shows an alignment of the Omp100 amino acid sequence with the most similar sequence found in a search of the GenBank database.

Figure 5 shows an alignment of the PonA amino acid sequence with the most similar sequence found in a search of the GenBank database.

Figure 6 shows an alignment of the HtrA amino acid sequence with the most similar sequence found in a search of the GenBank database.

Figure 7 shows an alignment of the HypC amino acid sequence with the most similar sequence found in a search of the GenBank database.

Figure 8 shows an alignment of the Orf1 amino acid sequence with the most similar sequence found in a search of the GenBank database.

Figure 9 shows an alignment of the LysS amino acid sequence with the most similar sequence found in a search of the GenBank database.

DETAILED DESCRIPTION OF THE INVENTION

[0018] All patents, patent applications, and publications cited herein are hereby incorporated by reference in their entireties.

Polynucleotide Molecules

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[0019] An isolated polynucleotide molecule of the present invention can have a nucleotide sequence derived from any species or strain of *Lawsonia*, but is preferably from the species *intracellularis*. Pathogenic strains or species of *Lawsonia* for use in practicing the present invention can be isolated from organs, tissues or body fluids of infected animals using isolation techniques as described below.

[0020] As used herein, the terms "polynucleotide molecule," "polynucleotide sequence," "coding sequence," "open-reading frame (ORF)," and the like, are intended to refer to both DNA and RNA molecules, which can either be single-stranded or double-stranded, and that can include one or more prokaryotic sequences, cDNA sequences, genomic DNA sequences including exons and introns, and chemically synthesized DNA and RNA sequences, and both sense and corresponding anti-sense strands. As used herein, the term "ORF" refers to the minimal nucleotide sequence required to encode a *Lawsonia* HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein, without any intervening termination codons.

[0021] Production and manipulation of the polynucleotide molecules and oligonucleotide molecules disclosed herein are within the skill in the art and can be carried out according to recombinant techniques described, among other places, in Maniatis et al., 1989, Molecular Cloning. A Laboratory Manual, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY; Ausubel et al., 1989, Current Protocols in Molecular Biology, Greene Publishing Associates & Wiley Interscience, NY; Sambrook et al., 1989, Molecular Cloning: A Laboratory Manual, 2d ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY; Innis et al. (eds), 1995, PCR Strategies, Academic Press, Inc., San Diego; and Erlich (ed), 1992, PCR Technology, Oxford University Press, New York and all revisions of these references.

[0022] References herein to the nucleotide sequences shown in SEQ ID NOS: 1 AND 2, and to substantial portions thereof, are intended to also refer to the corresponding nucleotide sequences and substantial portions thereof, respectively, as present in the following plasmids contained in *E. coli* Top10 cells deposited by Pfizer Inc. at Central Research, Eastern Point Road, Groton, CT, 06340 with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, MD, 20852:

pER432 containing the *ponA* gene and accorded ATCC accession number PTA-635, deposited on September 9, 1999;

pER434 containing the htrA gene and accorded ATCC accession number PTA-636, deposited on September 9, 1999;

pER436 containing the *hypC* gene and accorded ATCC accession number PTA-637, deposited on September 9, 1999;

pER438 containing the *ycfW* and *abc1* genes and accorded ATCC accession number PTA-638, deposited on September 9, 1999;

pER440 containing the *omp100* gene and accorded ATCC accession number PTA-639, deposited on September 9, 1999; and

pT068 containing the *lysS* and *ycfW* genes and accorded ATCC accession number PTA-2232, deposited on July 14, 2000.

40 [0023] In addition, references herein to the amino acid sequences shown in SEQ ID NOS:3-9, and SEQ ID NO: 102, and to substantial portions and peptide fragments thereof, are intended to also refer to the corresponding amino acid sequences, and substantial portions and peptide fragments thereof, respectively, encoded by the corresponding protein encoding nucleotide sequences present in the plasmids listed above, unless otherwise indicated.

45 HtrA-Related Polynucleotide Molecules

The present invention provides an isolated polynucleotide molecule comprising a nucleotide sequence encoding the HtrA protein from *L. intracellularis*. In a preferred embodiment, the HtrA protein has the amino acid sequence of **SEQ ID NO**: **7**. In a further preferred embodiment, the isolated HtrA-encoding polynucleotide molecule of the present invention comprises a nucleotide sequence selected from the group consisting of the nucleotide sequence of **SEQ ID NO**: **2** from about nt 2891 to about nt 4315, which is the nucleotide sequence of the open reading frame (ORF) of the *htrA* gene, and the nucleotide sequence of the HtrA-encoding ORF of plasmid pER434 (ATCC accession number PTA-636).

[0025] The present invention further provides an isolated polynucleotide molecule having a nucleotide sequence that is homologous to the nucleotide sequence of a HtrA-encoding polynucleotide molecule of the present invention. The term "homologous" when used to refer to a HtrA-related polynucleotide molecule means a polynucleotide molecule having a nucleotide sequence: (a) that encodes the same protein as one of the aforementioned HtrA-encoding polynucleotide molecules of the present invention, but that includes one or more silent changes to the nucleotide sequence

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according to the degeneracy of the genetic code; or (b) that hybridizes to the complement of a polynucleotide molecule having a nucleotide sequence that encodes the amino acid sequence of the *L. intracellularis* HtrA protein under at least moderately stringent conditions, *i.e.*, hybridization to filter-bound DNA in 0.5 M NaHPO₄, 7% sodium dodecyl sulfate (SDS), 1 mM EDTA at 65°C, and washing in 0.2xSSC/0.1% SDS at 42°C (see Ausubel *et al.* (eds.), 1989, Current Protocols in Molecular Biology, Vol. I, Green Publishing Associates, Inc., and John Wiley & Sons, Inc., New York, at p. 2.10.3), and that is useful in practicing the present invention. In a preferred embodiment, the homologous polynucleotide molecule hybridizes to the complement of a polynucleotide molecule having a nucleotide sequence that encodes the amino acid sequence of the *L. intracellularis* HtrA protein under highly stringent conditions, *i.e.*, hybridization to filter-bound DNA in 0.5 M NaHPO₄, 7% SDS, 1 mM EDTA at 65°C, and washing in 0.1xSSC/0.1% SDS at 68°C (Ausubel *et al.*, 1989, above), and is useful in practicing the present invention. In a more preferred embodiment, the homologous polynucleotide molecule hybridizes under highly stringent conditions to the complement of a polynucleotide molecule consisting of a nucleotide sequence selected from the group consisting of the HtrA encoding ORF of SEQ ID NO: 2, which is from about nt 2891 to about nt 4315. As noted above, reference to homologous polynucleotide molecules

[0026] As used herein, a polynucleotide molecule is "useful in practicing the present invention" where the polynucleotide molecule can be used to amplify a Lawsonia-specific polynucleotide molecule using a standard amplification technique, such as the polymerase chain reaction, or as a diagnostic reagent to detect the presence of a Lawsonia-specific polynucleotide in a fluid or tissue sample from a Lawsonia-infected animal, or where the polynucleotide molecule encodes a polypeptide that is useful in practicing the invention, as described below.

[0027] Polynucleotide molecules of the present invention having nucleotide sequences that are homologous to the nucleotide sequence of a HtrA-encoding polynucleotide molecule of the present invention do not include polynucleotide molecules that have been described from bacteria such as *E. coli*, *S. typhimurium*, *C. jejuni*, *H. influenzae*, *B. melitensis*, *B. abortus*, *C. trachomatis*, *Y. enterocolitia*, *Rickettsia*, *B. burgdorferi*, and *B. subtilis*. The *L. intracellularis* HtrA protein encoded by SEQ ID NO: 2 has 39.6% identity of amino acid sequence with the *B. abortus* HtrA protein. The *L. intracellularis* protein is 474 residues in length and the *B. abortus* protein is 513 residues in length. The *L. intracellularis* protein is 35.4% identical to that of *H. influenzae*.

[0028] The homologous nucleotide sequence of the molecule of the invention preferably comprises a sequence that has more than 50%, more preferably more than about 90%, even more preferably more than about 95%, and most preferably more than about 99% sequence identity to the molecule of SEQ ID NO: 2, which is from about nt 2891 to about nt 4315, wherein sequence identity is determined by use of the BLASTN algorithm (GenBank, National Center for Biotechnology Information).

[0029] In another embodiment, the polynucleotide has a homologous sequence that is more than about 50% of the length of the nucleotide sequence encoding the *L. intracellularis* HtrA protein. In another embodiment the sequence is more than 70%, in another embodiment the sequence is more than 90%, and in another embodiment more than about 98%, of the length of the nucleotide sequence encoding the *L. intracellularis* protein. In yet another embodiment, the isolated polynucleotide that has a homologous sequence is equal in length to the sequence encoding the *L. intracellularis* HtrA protein.

[0030] In yet another embodiment, the nucleotide sequence that is homologous to the *L. intracellularis* HtrA protein encoding sequence has between 1 and 50, more preferably between 1 and 25, and most preferably between 1 and 5 nucleotides inserted, deleted, or substituted with respect to the sequence of **SEQ ID NO**: 2 which is from about nt 2891 to about nt 4315.

The present invention further provides an isolated polynucleotide molecule comprising a nucleotide [0031] sequence that encodes a polypeptide that is homologous to the L. intracellularis HtrA protein. As used herein to refer to polypeptides that are homologous to the L. intracellularis HtrA protein, the term "homologous" refers to a polypeptide otherwise having the amino acid sequence of the L. intracellularis HtrA protein, but in residues has been substituted with a different amino acid residue, where the resulting polypeptide is useful in practicing the present invention. Conservative amino acid substitutions are well-known in the art. tions include those described by Dayhof, M.D., 1978, Nat. Biomed. Res. Found., Washington, D.C., Vol. 5, Sup. 3, among others. More specifically, conservative amino acid substitutions are those that generally take place within a familia or an interest of the second substitutions. ily of amino acids that are related in acidity, polarity, or bulkiness of their side chains. Genetically encoded amino acids are generally divided into four groups: (1) acidic = aspartate, glutamate; (2) basic = lysine, arginine, histidine; (3) nonpolar = alanine, valine, leucine, isoleucine, proline, phenytalanine, methionine, tryptop han; and (4) uncharged polar = glycine, asparagine, glutamine, cysteine, serine, threonine, and tyrosine. Phenylalan ine, tryptophan and tyrosine are also jointly classified as aromatic amino acids. One or more replacements within any particular group, e.g., of a leucine with an isoleucine or valine, or of an aspartate with a glutamate, or of a threonine with acid residue with a structurally related amino acid residue, e.g., an amino acid residue with similar acidity, polarity, bulkiness of side chain, or with similarity in some combination thereof, will generally have an insignificant effect on the function or immunogenicity of the polypeptide. In a preferred embodiment, the homologous polypeptide has at least about 50%, more preferably at least about 70%, and even more preferably at least about 90% sequence identity, and most preferably at least 95% sequence identity to SEQ ID NO: 7.

[0032] In another embodiment, the polynucleotide encodes an isolated polypeptide consisting of the *L. intracellularis* HtrA protein having between 1 and 10, and more preferably between 1 and 5, amino acids inserted, deleted, or substituted, including combinations thereof. In a more particular example of this embodiment, the polynucle-otide encodes an isolated polypeptide having between 1 and 5 amino acids conservatively substituted for the HtrA sequence of SEO ID NO: 7.

[0033] As used herein, a polypeptide is "useful in practicing the present invention" where the polypeptide can be used as a diagnostic reagent to detect the presence of *Lawsonia*-specific antibodies in a blood, serum, or other biologif it can be used to induce an immune response in an animal against *Lawsonia*. The polypeptide is also useful [0034] The present invention further provides a party of the present invention further provides a party.

[0034] The present invention further provides a polynucleotide molecule consisting of a substantial portion of any of the aforementioned *Lawsonia HtrA*-related polynucleotide molecules of the present invention. As used herein, a "substantial portion" of a *HtrA*-related polynucleotide molecule means a polynucleotide molecule consisting of less than the complete nucleotide sequence of the *HtrA*-related polynucleotide molecule, but comprising at least about 5%, more preferably at least about 10%, and even more preferably at least about 20%, and most preferably at least about 50% of the nucleotide sequence of the *HtrA*-related polynucleotide molecule, and that is useful in practicing the present invention. Such polynucleotide molecules include, for example polynucleotide molecules encoding peptide fragments of the HtrA protein.

20 [0035] In addition to the nucleotide sequences of any of the aforementioned HtrA-related polynucleotide molecules, polynucleotide molecules of the present invention can further comprise, or alternatively may consist of, nucleotide sequences selected from those that naturally flank the HtrA ORF or gene in situ in L. intracellularis, and include the nucleotide sequences shown in SEQ ID NO: 2 from about nt 2691 to about nt 2890 and from about nt 4316 to about nt 4580, or substantial portions thereof.

PonA -Related Polynucleotide Molecules

[0036] The present invention provides an isolated polynucleotide molecule comprising a nucleotide sequence encoding the PonA protein from *L. intracellularis*. In a preferred embodiment, the PonA protein has the amino acid sequence of SEQ ID NO: 6. In a further preferred embodiment, the isolated PonA-encoding polynucleotide molecule of the present invention comprises a nucleotide sequence selected from the group consisting of the nucleotide sequence of SEQ ID NO: 2 from about nt 252 to about nt 2690 (the nucleotide sequence of the open reading frame (ORF) of the *PonA* gene) and the nucleotide sequence of the PonA-encoding ORF of plasmid pER432 (ATCC accession number PTA-635).

[0037] The present invention further provides an isolated polynucleotide molecule having a nucleotide sequence that is "homologous" to the nucleotide sequence of a PonA-encoding polynucleotide molecule of the present invention, as that term is correspondingly defined above with respect to HtrA related polynucleotide molecules. In a preferred embodiment, the homologous polynucleotide molecule hybridizes to the complement of a polynucleotide molecule having a nucleotide sequence that encodes the amino acid sequence of the *L. intracellularis* PonA protein under highly stringent conditions. In a more preferred embodiment, the homologous polynucleotide molecule hybridizes under highly stringent conditions to the complement of a polynucleotide molecule consisting of a nucleotide sequence of SEQ ID NO: 2 from about nt 252 to about nt 2690.

[0038] Polynucleotide molecules of the present invention having nucleotide sequences that are homologous to the nucleotide sequence of a PonA-encoding polynucleotide molecule of the present invention do not include known polynucleotide molecules encoding PonA proteins of *Neisseria flavescens*, *N. gonorrhoeae*, and *N. meningitidis*.

[0039] The homologous nucleotide sequence of the molecule of the invention preferably comprises a sequence that has more than 50%, more preferably more than about 90%, even more preferably more than about 95%, and most preferably more than about 99% sequence identity to the molecule of SEQ ID NO: 2, which is from about nt 252 to about technology information).

[0040] In another embodiment, the polynylactical is

[0040] In another embodiment, the polynucleotide has a homologous sequence that is more than about 50% of the length of the nucleotide sequence encoding the *L. intracellularis* PonA protein. In another embodiment, the sequence than 90%, and in another embodiment more than about 98%, of the length of the nucleotide sequence encoding the *L. intracellularis* protein. In yet another embodiment, the isolated polynucleotide that has a homologous sequence is equal in length to the sequence encoding the *L. intracellularis* PonA protein.

[0041] In yet another embodiment, the nucleotide sequence that is homologous to the *L. intracellularis* PonA protein encoding sequence has between 1 and 50, more preferably between 1 and 25, and most preferably between 1 and 5 nucleotides inserted, deleted, or substituted with respect to the sequence of SEQ ID NO: 2.

[0042] The present invention further provides an isolated polynucleotide molecule comprising a nucleotide sequence that encodes a polypeptide that is "homologous" to the *L. intracellularis* PonA protein, as that term is correspondingly described with respect to the HtrA protein above. In a preferred embodiment, the homologous polypeptide has at least about 50%, more preferably at least about 70%, and even more preferably at least about 90% sequence identity, and most preferably at least 95% sequence identity to SEQ ID NO: 6.

[0043] In another embodiment, the polynucleotide encodes an isolated polypeptide consisting of the *L. intracellularis* PonA protein having between 1 and 10, and more preferably between 1 and 5, amino acids inserted, deleted, or substituted, including combinations thereof. In a more particular example of this embodiment, the polynucleotide encodes an isolated polypeptide having between 1 and 5 amino acids conservatively substituted for the PonA sequence of SEQ ID NO: 6.

[0044] The present invention further provides a polynucleotide molecule consisting of a "substantial portion" of any of the aforementioned *Lawsonia-PonA-*related polynucleotide molecules of the present invention, as that term is correspondingly described above with respect to the HtrA protein.

[0045] In addition to the nucleotide sequences of any of the aforementioned *PonA*-related polynucleotide molecules, polynucleotide molecules of the present invention can further comprise, or alternatively may consist of, nucleotide sequences selected from those that naturally flank the *ponA* ORF or gene *in situ* in *L. intracellularis*, and include the nucleotide sequences shown in **SEQ ID NO: 2** from about nt 126 to about nt 251 and from about nt 2691 to about nt 2890, or substantial portions thereof.

HypC -Related Polynucleotide Molecules

[0046] The present invention provides an isolated polynucleotide molecule comprising a nucleotide sequence encoding the HypC protein from *L. intracellularis*. In a preferred embodiment, the HypC protein has the amino acid sequence of SEQ ID NO: 8. In a further preferred embodiment, the isolated HypC-encoding polynucleotide molecule of the present invention comprises a nucleotide sequence selected from the group consisting of the nucleotide sequence of SEQ ID NO: 2 from about nt 4581 to about nt 4829, and the nucleotide sequence of the HypC-encoding ORF of plasmid pER436 (ATCC accession number PTA-637).

[0047] The present invention further provides an isolated polynucleotide molecule having a nucleotide sequence that is "homologous" to the nucleotide sequence of a HypC-encoding polynucleotide molecule of the present invention, as that term is correspondingly defined above with respect to HtrA related polynucleotide molecules. In a preferred embodiment, the homologous polynucleotide molecule hybridizes to the complement of a polynucleotide molecule having a nucleotide sequence that encodes the amino acid sequence of the *L. intracellularis* HypC protein under highly stringent conditions. In a more preferred embodiment, the homologous polynucleotide molecule hybridizes under highly stringent conditions to the complement of a polynucleotide molecule consisting of a nucleotide sequence selected from the group consisting of the ORF of SEQ ID NO: 2 from about nt 4581 to about nt 4829

[0048] Polynucleotide molecules of the present invention having nucleotide sequences that are homologous to the nucleotide sequence of a HypC-encoding polynucleotide molecule of the present invention do not include polynucleotide molecules encoding HypC or HypD proteins of *Desulfovibrio gigas* and *Rizobium leguminosarum*.

[0049] The homologous nucleotide sequence of the molecule of the invention preferably comprises a sequence that has more than 50%, more preferably more than about 90%, even more preferably more than about 95%, and most preferably more than about 99% sequence identity to the molecule of SEQ ID NO: 2, which is from about nt 4581 to about nt 4829, wherein sequence identity is determined by use of the BLASTN algorithm (GenBank, National Center for Biotechnology Information).

[0050] In another embodiment, the polynucleotide has a homologous sequence that is more than about 50% of the length of the nucleotide sequence encoding the *L. intracellularis* HypC protein. In another embodiment, the sequence is more than 90%, and in another embodiment more than about 98%, of the length of the nucleotide sequence encoding the *L. intracellularis* HypC protein. In yet another embodiment, the isolated polynucleotide that has a homologous sequence is equal in length to the sequence encoding the *L. intracellularis* HypC protein.

[0051] In yet another embodiment, the nucleotide sequence that is homologous to the *L. intracellularis* HypC protein encoding sequence has between 1 and 50, more preferably between 1 and 25, and most preferably between 1 and 5 nucleotides inserted, deleted, or substituted with respect to the sequence of SEQ ID NO: 2.

[0052] The present invention further provides an isolated polynucleotide molecule comprising a nucleotide sequence that encodes a polypeptide that is "homologous" to the *L. intracellularis* HypC protein, as that term is correspondingly described with respect to the HtrA protein above. In a preferred embodiment, the homologous polypeptide has at least about 50%, more preferably at least about 70%, and even more preferably at least about 90% sequence identity, and most preferably at least 95% sequence identity to SEQ ID NO: 8.

[0053] In another embodiment, the polynucleotide encodes an isolated polypeptide consisting of the L. intracellularis HypC protein having between 1 and 10, and more preferably between 1 and 5, amino acids inserted,

deleted, or substituted, including combinations thereof. In a more particular example of this embodiment, the polynucleotide encodes an isolated polypeptide having between 1 and 5 amino acids conservatively substituted for the HypC sequence of SEQ ID NO: 8.

[0054] The present invention further provides a polynucleotide molecule consisting of a "substantial portion" of any of the aforementioned *Lawsonia HypC*-related polynucleotide molecules of the present invention, as that term is correspondingly described above with respect to the HtrA protein.

[0055] In addition to the nucleotide sequences of any of the aforementioned *HypC*-related polynucleotide molecules, polynucleotide molecules of the present invention can further comprise, or alternatively may consist of, nucleotide sequences selected from those that naturally flank the *hypC* ORF or gene *in situ* in *L. intracellularis*, and include the nucleotide sequences shown in **SEQ ID NO: 2** from about nt 4316 to about nt 4580 and from about nt 4830 to about nt 4911, or substantial portions thereof.

LysS -Related Polynucleotide Molecules

The present invention provides an isolated polynucleotide molecule comprising a nucleotide sequence encoding a LysS protein from *L. intracellularis*. In a preferred embodiment, the LysS protein has the amino acid sequence of **SEQ ID NO: 102**. In a further preferred embodiment, the isolated LysS-encoding polynucleotide molecule of the present invention comprises a nucleotide sequence selected from the group consisting of the nucleotide sequence of **SEQ ID NO: 1** from about nt 165 to about nt 1745 of the nucleotide sequence of the *lysS* gene, and the nucleotide sequence of the *LysS*-encoding ORF of plasmid pT068 (ATCC accession number PTA-2232).

The present invention further provides an isolated polynucleotide molecule having a nucleotide sequence that is "homologous" to the nucleotide sequence of a LysS-encoding polynucleotide molecule of the present invention, as that term is correspondingly defined above with respect to HtrA related polynucleotide molecules. In a preferred embodiment, the homologous polynucleotide molecule hybridizes to the complement of a polynucleotide molecule having a nucleotide sequence that encodes the amino acid sequence of the *L. intracellularis* LysS protein under highly stringent conditions. In a more preferred embodiment, the homologous polynucleotide molecule hybridizes under highly stringent conditions to the complement of a polynucleotide molecule consisting of a nucleotide sequence of SEQ ID NO: 1 from about nt 165 to about nt 1745.

[0058] The homologous nucleotide sequence of the molecule of the invention preferably comprises a sequence that has more than 50%, more preferably more than about 90%, even more preferably more than about 95%, and most preferably more than about 99% sequence identity to the molecule of SEQ ID NO: 1 from about nt 165 to about nt 1745, wherein sequence identity is determined by use of the BLASTN algorithm (GenBank, National Center for Biotechnology Information).

[0059] In another embodiment, the polynucleotide has a homologous sequence that is more than about 50% of the length of the nucleotide sequence encoding the *L. intracellularis* LysS protein. In another embodiment, the sequence is more than 90%, and in another embodiment more than about 98%, of the length of the nucleotide sequence encoding the *L. intracellularis* LysS protein. In yet another embodiment, the isolated polynucleotide that has a homologous sequence is equal in length to the sequence encoding the *L. intracellularis* LysS protein.

[0060] In yet another embodiment, the nucleotide sequence that is homologous to the *L. intracellularis* LysS protein encoding sequence has between 1 and 50, more preferably between 1 and 25, and most preferably between 1 and 5 nucleotides inserted, deleted, or substituted with respect to the sequence of SEQ ID NO: 1.

[0061] The present invention further provides an isolated polynucleotide molecule comprising a nucleotide sequence that encodes a polypeptide that is "homologous" to the *L. intracellularis* LysS protein, as that term is correspondingly described with respect to the HtrA protein above. In a preferred embodiment, the homologous polypeptide has at least about 50%, more preferably at least about 70%, and even more preferably at least about 90% sequence identity, and most preferably at least 95% sequence identity to SEQ ID NO: 102.

[0062] In another embodiment, the polynucleotide encodes an isolated polypeptide consisting of the *L. intracellularis* LysS protein having between 1 and 10, and more preferably between 1 and 5, amino acids inserted, deleted, or substituted, including combinations thereof. In a more particular example of this embodiment, the polynucleotide encodes an isolated polypeptide having between 1 and 5 amino acids conservatively substituted for the LysS sequence of SEQ ID NO: 102.

[0063] The present invention further provides a polynucleotide molecule consisting of a "substantial portion" of any of the aforementioned *Lawsonia lysS*-related polynucleotide molecules of the present invention, as that term is correspondingly described above with respect to the HtrA protein.

In addition to the nucleotide sequences of any of the aforementioned lysS-related polynucleotide molecules, polynucleotide molecules of the present invention can further comprise, or alternatively may consist of, nucleotide sequences selected from those that naturally flank the lysS ORF or gene in situ in L. intracellularis.

YcfW -Related Polynucleotide Molecules

[0065] The present invention provides an isolated polynucleotide molecule comprising a nucleotide sequence encoding the YcfW protein from *L. intracellularis*. In a preferred embodiment, the YcfW protein has the amino acid sequence of SEQ ID NO: 3. In a further preferred embodiment, the isolated YcfW-encoding polynucleotide molecule of the present invention comprises a nucleotide sequence selected from the group consisting of the nucleotide sequence of SEQ ID NO: 1 from about nt 1745 to about nt 3028 of the nucleotide sequence of the YcfW gene, and the nucleotide sequence of the YcfW-encoding ORF of plasmids pER438 (ATCC accession number PTA-638) and pT068 (ATCC accession number PTA-2232). The present invention further provides an isolated polynucleotide molecule having a nucleotide sequence that is "homologous" to the nucleotide sequence of a YcfW-encoding polynucleotide molecule of the present invention, as that term is correspondingly defined above with respect to HtrA related polynucleotide molecules. In a preferred embodiment, the homologous polynucleotide molecule hybridizes to the complement of a polynucleotide molecule having a nucleotide sequence that encodes the amino acid sequence of the *L. intracellularis* YcfW protein under highly stringent conditions. In a more preferred embodiment, the homologous polynucleotide molecule hybridizes under highly stringent conditions to the complement of a polynucleotide molecule consisting of a nucleotide sequence of SEQ ID NO: 1 from about nt 1745 to about nt 3028.

[0066] The homologous nucleotide sequence of the molecule of the invention preferably comprises a sequence that has more than 50%, more preferably more than about 90%, even more preferably more than about 95%, and most preferably more than about 99% sequence identity to the molecule of SEQ ID NO: 1 from about nt 1745 to about nt 3028, wherein sequence identity is determined by use of the BLASTN algorithm (GenBank, National Center for Biotechnology Information).

[0067] In another embodiment, the polynucleotide has a homologous sequence that is more than about 50% of the length of the nucleotide sequence encoding the *L. intracellularis* YcfW protein. In another embodiment, the sequence is more than 90%, and in another embodiment more than about 98%, of the length of the nucleotide sequence encoding the *L. intracellularis* YcfW protein. In yet another embodiment, the isolated polynucleotide that has a homologous sequence is equal in length to the sequence encoding the *L. intracellularis* YcfW protein.

[0068] In yet another embodiment, the nucleotide sequence that is homologous to the *L. intracellularis* YcfW protein encoding sequence has between 1 and 50, more preferably between 1 and 25, and most preferably between 1 and 5 nucleotides inserted, deleted, or substituted with respect to the sequence of SEQ ID NO: 1.

[0069] The present invention further provides an isolated polynucleotide molecule comprising a nucleotide sequence that encodes a polypeptide that is "homologous" to the *L. intracellularis* YcfW protein, as that term is correspondingly described with respect to the HtrA protein above. In a preferred embodiment, the homologous polypeptide has at least about 50%, more preferably at least about 70%, and even more preferably at least about 90% sequence identity, and most preferably at least 95% sequence identity to SEQ ID NO: 3.

[0070] In another embodiment, the polynucleotide encodes an isolated polypeptide consisting of the L. intracellularis YcfW protein having between 1 and 10, and more preferably between 1 and 5, amino acids inserted, deleted, or substituted, including combinations thereof. In a more particular example of this embodiment, the polynucle-otide encodes an isolated polypeptide having between 1 and 5 amino acids conservatively substituted for the YcfW sequence of SEQ ID NO: 3.

[0071] The present invention further provides a polynucleotide molecule consisting of a "substantial portion" of any of the aforementioned *Lawsonia YcfW*-related polynucleotide molecules of the present invention, as that term is correspondingly described above with respect to the HtrA protein.

[0072] In addition to the nucleotide sequences of any of the aforementioned YcfW-related polynucleotide molecules, polynucleotide molecules of the present invention can further comprise, or alternatively may consist of, nucleotide sequences selected from those that naturally flank the ycfW ORF or gene in situ in L. intracellularis.

ABC1 -Related Polynucleotide Molecules

[0073] The present invention provides an isolated polynucleotide molecule comprising a nucleotide sequence encoding the ABC1 protein from *L. intracellularis*. In a preferred embodiment, the ABC1 protein has the amino acid sequence of SEQ ID NO: 4. In a further preferred embodiment, the isolated ABC1-encoding polynucleotide molecule of the present invention comprises a nucleotide sequence selected from the group consisting of the nucleotide sequence of SEQ ID NO: 1 from about nt 3031 to about nt 3738 (the nucleotide sequence of the open reading frame (ORF) of the *ABC1* gene) and the nucleotide sequence of the ABC1-encoding ORF of plasmid pER438 (ATCC accession number PTA-638).

[0074] The present invention further provides an isolated polynucleotide molecule having a nucleotide sequence that is "homologous" to the nucleotide sequence of an ABC1-encoding polynucleotide molecule of the present invention, as that term is correspondingly defined above with respect to HtrA related polynucleotide molecules. In a preferred

embodiment, the homologous polynucleotide molecule hybridizes to the complement of a polynucleotide molecule having a nucleotide sequence that encodes the amino acid sequence of the *L. intracellularis* ABC1 protein under highly stringent conditions. In a more preferred embodiment, the homologous polynucleotide molecule hybridizes under highly stringent conditions to the complement of a polynucleotide molecule consisting of a nucleotide sequence selected from the group consisting of the ORE of SEQ ID NO: 1, which is from about nt 3031 to about nt 3738.

[0075] Polynucleotide molecules of the present invention having nucleotide sequences that are homologous to the nucleotide sequence of a ABC1-encoding polynucleotide molecule of the present invention do not include polynucleotide molecules encoding ABC1 proteins of *Neisseria flavescens*, *N. gonorrhoeae*, and *N. meningitidis*.

[0076] The nucleotide sequence of the molecule of the invention preferably comprises a sequence that has more than 50%, more preferably more than about 90%, even more preferably more than about 95%, and most preferably more than about 99% sequence identity to the molecule of SEQ ID NO: 1 from about nt 3031 to about nt 3738, wherein sequence identity is determined by use of the BLASTN algorithm (GenBank, National Center for Biotechnology Information).

[0077] In another embodiment, the polynucleotide has a homologous sequence that is more than about 50% of the length of the nucleotide sequence encoding the *L. intracellularis* ABC1 protein. In another embodiment, the sequence is more than 90%, and in another embodiment more than about 98%, of the length of the nucleotide sequence encoding the *L. intracellularis* ABC1 protein. In yet another embodiment, the isolated polynucleotide that has a homologous sequence is equal in length to the sequence encoding the *L. intracellularis* ABC1 protein.

[0078] In yet another embodiment, the nucleotide sequence that is homologous to the *L. intracellularis* ABC1 protein encoding sequence has between 1 and 50, more preferably between 1 and 25, and most preferably between 1 and 5 nucleotides inserted, deleted, or substituted with respect to the sequence of SEQ ID NO: 1.

[0079] The present invention further provides an isolated polynucleotide molecule comprising a nucleotide sequence that encodes a polypeptide that is "homologous" to the *L. intracellularis* ABC1 protein, as that term is correspondingly described with respect to the HtrA protein above. In a preferred embodiment, the homologous polypeptide has at least about 50%, more preferably at least about 70%, and even more preferably at least about 90% sequence identity, and most preferably at least 95% sequence identity to SEQ ID NO: 1.

[0080] In another embodiment, the polynucleotide encodes an isolated polypeptide consisting of the *L. intracellularis* ABC1 protein having between 1 and 10, and more preferably between 1 and 5, amino acids inserted, deleted, or substituted, including combinations thereof. In a more particular example of this embodiment, the polynucleotide encodes an isolated polypeptide having between 1 and 5 amino acids conservatively substituted for the ABC1 sequence of SEQ ID NO: 4.

[0081] The present invention further provides a polynucleotide molecule consisting of a "substantial portion" of any of the aforementioned *Lawsonia ABC1*-related polynucleotide molecules of the present invention, as that term is correspondingly described above with respect to the HtrA protein.

[0082] In addition to the nucleotide sequences of any of the aforementioned ABC1-related polynucleotide molecules, polynucleotide molecules of the present invention can further comprise, or alternatively may consist of, nucleotide sequences selected from those that naturally flank the abc1 ORF or gene in situ in L. intracellularis, and include the flanking nucleotide sequences shown in SEQ ID NO: 1.

40 Omp100 -Related Polynucleotide Molecules

[0083] The present invention provides an isolated polynucleotide molecule comprising a nucleotide sequence encoding the Omp100 protein from *L. intracellularis*. In a preferred embodiment, the Omp100 protein has the amino acid sequence of SEQ ID NO: 5. In a further preferred embodiment, the isolated Omp100-encoding polynucleotide molecule of the present invention comprises a nucleotide sequence selected from the group consisting of the nucleotide sequence of SEQ ID NO: 1 from about nt 3695 to about nt 6385 (the nucleotide sequence of the open reading frame (ORF) of the *Omp100* gene), and the nucleotide sequence of the Omp100-encoding ORF of plasmid pER440 (ATCC accession number PTA-639).

[0084] The present invention further provides an isolated polynucleotide molecule having a nucleotide sequence that is "homologous" to the nucleotide sequence of a Omp100-encoding polynucleotide molecule of the present invention, as that term is correspondingly defined above with respect to HtrA related polynucleotide molecules. In a preferred embodiment, the homologous polynucleotide molecule hybridizes to the complement of a polynucleotide molecule having a nucleotide sequence that encodes the amino acid sequence of the *L intracellularis* Omp100 protein under highly stringent conditions. In a more preferred embodiment, the homologous polynucleotide molecule hybridizes under highly stringent conditions to the complement of a polynucleotide molecule consisting of a nucleotide sequence selected from the group consisting of the ORF of SEQ ID NO: 1, which is from about nt 3695 to about nt 6385.

[0085] Polynucleotide molecules of the present invention having nucleotide sequences that are homologous to the nucleotide sequence of a Omp100-encoding polynucleotide molecule of the present invention do not include polynucle-

otide molecules encoding any of the following proteins listed in the GenBank database: YaeT (Accn. U70214 or AE000127) of *E. coli*; Oma90 (Accn. AF120927) of *Shigella flexneri*, Omp85 (Accn. AF021245) of *Neisseria meningitidis*, D15 (Accn. U60834) of *Haemophilus influenzae* (D15), and Oma87 (Accn. U60439) of *Pasteurella multocida*.

[0086] The nucleotide sequence of the molecule of the invention preferably comprises a homologous sequence that has more than 50%, more preferably more than about 90%, even more preferably more than about 95%, and most preferably more than about 99% sequence identity to the molecule of SEQ ID NO: 1 from about nt 3695 to about nt 6385, wherein sequence identity is determined by use of the BLASTN algorithm (GenBank, National Center for Biotechnology Information).

[0087] In another embodiment, the polynucleotide has a homologous sequence that is more than about 50% of the length of the nucleotide sequence encoding the *L. intracellularis* Omp100 protein. In another embodiment, the sequence is more than 90%, and in another embodiment more than about 98%, of the length of the nucleotide sequence encoding the *L. intracellularis* Omp100 protein. In yet another embodiment, the isolated polynucleotide that has a homologous sequence is equal in length to the sequence encoding the *L. intracellularis* Omp100 protein.

[0088] In yet another embodiment, the nucleotide sequence that is homologous to the *L. intracellularis* Omp100 protein encoding sequence has between 1 and 50, more preferably between 1 and 25, and most preferably between 1 and 5 nucleotides inserted, deleted, or substituted with respect to the sequence of **SEQ ID NO: 5**.

[0089] The present invention further provides an isolated polynucleotide molecule comprising a nucleotide sequence that encodes a polypeptide that is "homologous" to the *L. intracellularis* Omp100 protein, as that term is correspondingly described with respect to the HtrA protein above. In a preferred embodiment, the homologous polypeptide has at least about 50%, more preferably at least about 70%, and even more preferably at least about 90% sequence identity, and most preferably at least 95% sequence identity to SEQ ID NO: 5.

[0090] In another embodiment, the polynucleotide encodes an isolated polypeptide consisting of the *L. intracellularis* Omp100 protein having between 1 and 10, and more preferably between 1 and 5, amino acids inserted, deleted, or substituted, including combinations thereof. In a more particular example of this embodiment, the polynucleotide encodes an isolated polypeptide having between 1 and 5 amino acids conservatively substituted for the Omp100 sequence of SEQ ID NO: 5.

[0091] The present invention further provides a polynucleotide molecule consisting of a "substantial portion" of any of the aforementioned *Lawsonia Omp100*-related polynucleotide molecules of the present invention, as that term is correspondingly described above with respect to the HtrA protein.

[0092] In addition to the nucleotide sequences of any of the aforementioned *Omp100*-related polynucleotide molecules, polynucleotide molecules of the present invention can further comprise, or alternatively may consist of, nucleotide sequences selected from those that naturally flank the *omp100* ORF or gene *in situ* in *L. intracellularis*, and include the nucleotide sequences shown in **SEQ ID NO: 1**.

Promoter Sequences

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[0093] The present invention also relates to a polynucleotide molecule comprising a nucleotide sequence greater than 20 nucleotides having promoter activity and found within SEQ ID NO: 2 from about nt 2691 to about nt 2890, or its complement. As further discussed below in the Examples, it has been determined that this region of the *Lawsonia* sequence contains a temperature responsive promoter for the *htrA* gene. In a preferred embodiment, the polynucleotide comprises the sequence of about nt 2797 to nt 2829.

[0094] The present invention also relates to oligonucleotides having promoter activity that hybridize under moderately stringent, and more preferably under highly stringent conditions, to the complement of the nucleotide sequence greater than 20 nucleotides having promoter activity and found within SEQ ID NO: 2 from about nt 2691 to about nt 2890. Preferably the oligonucleotide having promoter activity hybridizes under moderately stringent or highly stringent conditions to the complement of the polynucleotide comprising the sequence from about nt 2797 to nt 2829. In another embodiment, the invention encompasses an oligonucleotide having promoter activity having between 1 and 25, and most preferably between 1 and 5 nucleotides inserted, deleted, or substituted with respect to the sequence of SEQ ID NO: 2 which is from about nt 2691 to about nt 2890.

[0095] The functional sequences having promoter activity of the present invention are useful for a variety of purposes including for controlling the recombinant expression of any of the genes of the present invention, or of other genes or coding sequences, in host cells of *L. intracellularis* or in host cells of any other species of *Lawsonia*, or in any other appropriate host cell. Such other genes or coding sequences can either be native or heterologous to the recombinant host cell. The promoter sequence can be fused to the particular gene or coding sequence using standard recombinant techniques as known in the art so that the promoter sequence is in operative association therewith, as "operative association" is defined below. By using the promoter, recombinant expression systems can, for example, be constructed and used to screen for compounds and transcriptional factors that can modulate the expression of the genes

of Lawsonia or other bacteria. In addition, such promoter constructs can be used to express heterologous polypeptides in Lawsonia, E. coli, or other appropriate host cells.

Oligonucleotide Molecules

[0096] The present invention further provides oligonucleotide molecules that hybridize to any one of the aforementioned polynucleotide molecules of the present invention, or that hybridize to a polynucleotide molecule having a nucleotide sequence that is the complement of any one of the aforementioned polynucleotide molecules of the present invention. Such oligonucleotide molecules are preferably at least about 10 nucleotides in length, and more preferably from about 15 to about 30 nucleotides in length, and hybridize to one or more of the aforementioned polynucleotide molecules under highly stringent conditions, *i.e.*, washing in 6xSSC/0.5% sodium pyrophosphate at about 37°C for ~14-base oligos, at about 48°C for ~17-base oligos, at about 55°C for ~20-base oligos, and at about 60°C for ~23-base oligos. Other hybridization conditions for longer oligonucleotide molecules of the present invention can be determined by the skilled artisan using standard techniques. In a preferred embodiment, an oligonucleotide molecules of the present invention is complementary to a portion of at least one of the aforementioned polynucleotide molecules of the present invention.

[0097] The oligonucleotide molecules of the present invention are useful for a variety of purposes, including as primers in amplification of a *Lawsonia*-specific polynucleotide molecule for use, *e.g.*, in differential disease diagnosis, or to act as antisense molecules useful in gene regulation. Suitably designed primers can also be used to detect the presence of *Lawsonia*-specific polynucleotide molecules in a sample of animal tissue or fluid, including brain tissue, lung tissue, intestinal tissue, placental tissue, blood, cerebrospinal fluid, feces, mucous, urine, amniotic fluid, *etc.* The oligonucleotide molecule specifically reacts with the Lawsonia organism; this is generally accomplished by employing a sequence of sufficient length. The production of a specific amplification product can support a diagnosis of *Lawsonia* infection, while lack of an amplified product can point to a lack of infection. Methods for conducting amplifications, such as the polymerase chain reaction (PCR), are described, among other places, in Innis *et al.* (eds), 1995, above; and Erlich (ed), 1992, above. Other amplification techniques known in the art, *e.g.*, the ligase chain reaction, can alternatively be used. The sequences of the polynucleotide molecules disclosed herein can also be used to design primers for use in isolating homologous genes from other species or strains of *Lawsonia* or other bacterial cells.

[0098] Specific though non-limiting embodiments of oligonucleotide molecules useful in practicing the present invention include oligonucleotide molecules selected from the group consisting of SEQ ID NOS: 10-101 and the complements thereof.

Recombinant Expression Systems Cloning And Expression Vectors

The present invention further encompasses methods and compositions for cloning and expressing any of the polynucleotide molecules of the present invention, including cloning vectors, expression vectors, transformed host cells comprising any of said vectors, and novel strains or cell lines derived therefrom. In a preferred embodiment, the present invention provides a recombinant vector comprising a polynucleotide molecule having a nucleotide sequence encoding the HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein of *L. intracellularis*. In specific embodiments, the present invention provides plasmid pER432 containing the *ponA* gene (ATCC accession number PTA-635), plasmid pER434 containing the *htrA* gene (ATCC accession number PTA-636), plasmid pER436 containing the *htrA* gene (ATCC accession number PTA-637), plasmid pTO68 containing the *lysS* and *ycfW* genes (ATCC accession number PTA-2232), plasmid pER438 containing the *ycfW* and *abc1* genes (ATCC accession number PTA-638), and plasmid pER440 containing the Omp100 gene (ATCC accession number PTA-639). The invention also encompasses recombinant vectors and transformed cells employed to obtain polypeptides of the invention.

[0100] Recombinant vectors of the present invention, particularly expression vectors, are preferably constructed so that the coding sequence for the polynucleotide molecule of the invention is in operative association with one or more regulatory elements necessary for transcription and translation of the coding sequence to produce a polypeptide. As used herein, the term "regulatory element" includes but is not limited to nucleotide sequences that encode inducible and non-inducible promoters, enhancers, operators, ribosome-binding sites, and other elements known in the art that serve to drive and/or regulate expression of polynucleotide coding sequences. Also, as used herein, the coding sequence is in "operative association" with one or more regulatory elements where the regulatory elements effectively regulate and allow for the transcription of the coding sequence or the translation of its mRNA, or both.

[0101] Methods are well-known in the art for constructing recombinant vectors containing particular coding sequences in operative association with appropriate regulatory elements, and these can be used to practice the present invention. These methods include *in vitro* recombinant techniques, synthetic techniques, and *in vivo* genetic recombination. See, *e.g.*, the techniques described in Maniatis *et al.*, 1989, above; Ausubel *et al.*, 1989, above; Sambrook *et al.*, 1989, above; Innis et al., 1995, above; and Erlich, 1992, above.

[0102] A variety of expression vectors are known in the art which can be utilized to express the HtrA, PonA, HypC. LysS, YcfW, ABC1, or Omp100 coding sequences of the present invention, including recombinant bacteriophage DNA, plasmid DNA, and cosmid DNA expression vectors containing the particular coding sequences. Typical prokaryotic expression vector plasmids that can be engineered to contain a polynucleotide molecule of the present invention include pUC8, pUC9, pBR322 and pBR329 (Biorad Laboratories, Richmond, CA), pPL and pKK223 (Pharmacia, Piscataway, NJ), pQE50 (Qiagen, Chatsworth, CA), and pGEM-T EASY (Promega, Madison, WI), among many others. Typical eukaryotic expression vectors that can be engineered to contain a polynucleotide molecule of the present invention include an ecdysone-inducible mammalian expression system (Invitrogen, Carlsbad, CA), cytomegalovirus promoter-enhancer-based systems (Promega, Madison, WI; Stratagene, La Jolla, CA, Invitrogen), and baculovirus-based expression systems (Promega), among others.

[0103] The regulatory elements of these and other vectors can vary in their strength and specificities. Depending on the host/vector system utilized, any of a number of suitable transcription and translation elements can be used. For instance, when cloning in mammalian cell systems, promoters isolated from the genome of mammalian cells, e.g., mouse metallothionein promoter, or from viruses that grow in these cells, e.g., vaccinia virus 7.5K promoter or Moloney murine sarcoma virus long terminal repeat, can be used. Promoters obtained by recombinant DNA or synthetic techniques can also be used to provide for transcription of the inserted sequence. In addition, expression from certain promoters can be elevated in the presence of particular inducers, e.g., zinc and cadmium ions for metallothionein promoters. Non-limiting examples of transcriptional regulatory regions or promoters include for bacteria, the β -gal promoter, the T7 promoter, the TAC promoter, λ left and right promoters, trp and lac promoters, trp-lac fusion promoters, etc.; for yeast, glycolytic enzyme promoters, such as ADH-I and -II promoters, GPK promoter, PGI promoter, TRP promoter, etc.; and for mammalian cells, SV40 early and late promoters, adenovirus major late promoters, among others. The present invention further provides a polynucleotide molecule comprising the nucleotide sequence of the promoter of the htrA gene of L. intracellularis, which can be used to express any of the coding sequences of the present invention in Lawsonia, E. coli, or other suitable hosts.

[0104] Specific initiation signals are also required for sufficient translation of inserted coding sequences. These signals typically include an ATG initiation codon and adjacent sequences. In cases where the polynucleotide molecule of the present invention including its own initiation codon and adjacent sequences are inserted into the appropriate expression vector, no additional translation control signals may be needed. However, in cases where only a portion of a coding sequence is inserted, exogenous translational control signals, including the ATG initiation codon, may be required. These exogenous translational control signals and initiation codons can be obtained from a variety of sources, both natural and synthetic. Furthermore, the initiation codon must be in phase with the reading frame of the coding regions to ensure in-frame translation of the entire insert.

[0105] Expression vectors can also be constructed that will express a fusion protein comprising a protein or polypeptide of the present invention. Such fusion proteins can be used, *e.g.*, to raise antisera against a *Lawsonia* protein, to study the biochemical properties of the *Lawsonia* protein, to engineer a *Lawsonia* protein exhibiting different immunological or functional properties, or to aid in the identification or purification, or to improve the stability, of a recombinantly-expressed *Lawsonia* protein. Possible fusion protein expression vectors include but are not limited to vectors incorporating sequences that encode β-galactosidase and trpE fusions, maltose-binding protein fusions (pMal series; New England Biolabs), glutathione-S-transferase fusions (pGEX series; Pharmacia), polyhistidine fusions (pET series; Novagen Inc., Madison, WI), and thioredoxin fusions (pTrxFus; Invitrogen, Carlsbad, CA). Methods are well-known in the art for constructing expression vectors encoding these and other fusion proteins.

[0106] The fusion protein can be useful to aid in purification of the expressed protein. In non-limiting embodiments, e.g., a HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100-maltose-binding fusion protein can be purified using amylose resin; a HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100-glutathione-S-transferase fusion protein can be purified using glutathione-agarose beads; and a HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100-polyhistidine fusion protein can be purified using divalent nickel resin. Alternatively, antibodies against a carrier protein or peptide can be used for affinity chromatography purification of the fusion protein. For example, a nucleotide sequence coding for the target epitope of a monoclonal antibody can be engineered into the expression vector in operative association with the regulatory elements and situated so that the expressed epitope is fused to a *Lawsonia* protein of the present invention. In a non-limiting embodiment, a nucleotide sequence coding for the FLAG™ epitope tag (International Biotechnologies Inc.), which is a hydrophilic marker peptide, can be inserted by standard techniques into the expression vector at a point corresponding to the amino or carboxyl terminus of the HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein. The expressed HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein-FLAG™ epitope fusion product can then be detected and affinity-purified using commercially available anti-FLAG™ antibodies.

[0107] The expression vector can also be engineered to contain polylinker sequences that encode specific protease cleavage sites so that the expressed *Lawsonia* protein can be released from the carrier region or fusion partner by treatment with a specific protease. For example, the fusion protein vector can include a nucleotide sequence encoding a thrombin or factor Xa cleavage site, among others.

[0108] A signal sequence upstream from and in the same reading frame with the *Lawsonia* coding sequence can be engineered into the expression vector by known methods to direct the trafficking and secretion of the expressed protein. Non-limiting examples of signal sequences include those from α -factor, immunoglobulins, outer membrane proteins, penicillinase, and T-cell receptors, among others.

[0109] To aid in the selection of host cells transformed or transfected with a recombinant vector of the present invention, the vector can be engineered to further comprise a coding sequence for a reporter gene product or other selectable marker. Such a coding sequence is preferably in operative association with the regulatory elements, as described above. Reporter genes that are useful in practicing the invention are well-known in the art and include those encoding chloramphenicol acetyltransferase (CAT), green fluorescent protein, firefly luciferase, and human growth hormone, among others. Nucleotide sequences encoding selectable markers are well-known in the art, and include those that encode gene products conferring resistance to antibiotics or anti-metabolites, or that supply an auxotrophic requirement. Examples of such sequences include those that encode thymidine kinase activity, or resistance to methotrexate, ampicillin, kanamycin, chloramphenicol, zeocin, pyrimethamine, aminoglycosides, or hygromycin, among others.

Transformation Of Host Cells

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[0110] The present invention further provides transformed host cells comprising a polynucleotide molecule or recombinant vector of the present invention, and cell lines derived therefrom. Host cells useful in practicing the invention can be eukaryotic or prokaryotic cells. Such transformed host cells include but are not limited to microorganisms, such as bacteria transformed with recombinant bacteriophage DNA, plasmid DNA or cosmid DNA vectors, or yeast transformed with a recombinant vector, or animal cells, such as insect cells infected with a recombinant virus vector, e.g., baculovirus, or mammalian cells infected with a recombinant virus vector, e.g., adenovirus or vaccinia virus, among others. For example, a strain of *E. coli* can be used, such as, e.g., the DH5α strain available from the ATCC, Rockville, MD, USA (Accession No. 31343), or from GIBCO BRL, Gaithersburg, MD. Eukaryotic host cells include yeast cells, although mammalian cells, e.g., from a mouse, hamster, cow, monkey, or human cell line, among others, can also be utilized effectively. Examples of eukaryotic host cells that can be used to express a recombinant protein of the invention include Chinese hamster ovary (CHO) cells (e.g., ATCC Accession No. CCL-61), NIH Swiss mouse embryo cells NIH/3T3 (e.g., ATCC Accession No. CRL-1658), and Madin-Darby bovine kidney (MDBK) cells (ATCC Accession No. CCL-22). Transfected cells can express the polynucleotide of the invention by chromosomal integration, or episomally.

[0111] The recombinant vector of the invention is preferably transformed or transfected into one or more host cells of a substantially homogeneous culture of cells. The vector is generally introduced into host cells in accordance with known techniques, such as, e.g., by protoplast transformation, calcium phosphate precipitation, calcium chloride treatment. microinjection, electroporation, transfection by contact with a recombined virus, liposome-mediated transfection, DEAE-dextran transfection, transduction, conjugation, or microprojectile bombardment, among others. Selection of transformants can be conducted by standard procedures, such as by selecting for cells expressing a selectable marker. e.g., antibiotic resistance, associated with the recombinant expression vector.

[0112] Once an expression vector is introduced into the host cell, the integration and maintenance of the polynucleotide molecule of the present invention, either in the host cell genome or episomally, can be confirmed by standard techniques, e.g., by Southern hybridization analysis, restriction enzyme analysis, PCR analysis including reverse
transcriptase PCR (rt-PCR), or by immunological assay to detect the expected protein product. Host cells containing
and/or expressing a polynucleotide molecule of the present invention can be identified by any of at least four general
approaches that are well-known in the art, including: (i) DNA-DNA, DNA-RNA, or RNA-antisense RNA hybridization; (ii)
detecting the presence of "marker" gene functions; (iii) assessing the level of transcription as measured by the expression of specific mRNA transcripts in the host cell; or (iv) detecting the presence of mature polypeptide product, e.g., by
immunoassay, as known in the art.

Expression And Purification Of Recombinant Polypeptides

[0113] Once a polynucleotide molecule of the present invention has been stably introduced into an appropriate host cell, the transformed host cell is clonally propagated, and the resulting cells are grown under conditions conducive to the maximum production of the encoded polypeptide. Such conditions typically include growing transformed cells to high density. Where the expression vector comprises an inducible promoter, appropriate induction conditions such as, e.g., temperature shift, exhaustion of nutrients, addition of gratuitous inducers (e.g., analogs of carbohydrates, such as isopropyl-β-D-thiogalactopyranoside (IPTG)), accumulation of excess metabolic by-products, or the like, are employed as needed to induce expression.

[0114] Where the polypeptide is retained inside the host cells, the cells are harvested and lysed, and the product is substantially purified or isolated from the lysate under extraction conditions known in the art to minimize protein degra-

dation such as, e.g., at 4°C, or in the presence of protease inhibitors, or both. Where the polypeptide is secreted from the host cells, the exhausted nutrient medium can simply be collected and the polypeptide substantially purified or isolated therefrom.

[0115] The polypeptide can be substantially purified or isolated from cell lysates or culture medium, as necessary, using standard methods, including but not limited to one or more of the following methods: ammonium sulfate precipitation, size fractionation, ion exchange chromatography, HPLC, density centrifugation, and affinity chromatography. If the polypeptide lacks biological activity, it can be detected as based, e.g., on size, or reactivity with a polypeptide-specific antibody, or by the presence of a fusion tag. For use in practicing the present invention, the polypeptide can be in an unpurified state as secreted into the culture fluid or as present in a cell lysate, but is preferably substantially purified or isolated therefrom. As used herein, a polypeptide is "substantially purified" where the polypeptide constitutes at least about 20 wt% of the protein in a particular preparation. Also, as used herein, a polypeptide is "isolated" where the polypeptide constitutes at least about 80 wt% of the protein in a particular preparation. In another embodiment of the invention, the protein is present in a preparation in at least about a 1000x higher concentration than its natural counterpart is normally found in a preparation of *L. intracellularis* cell lysate.

Polypeptides

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[0116] Thus, the present invention encompasses a substantially purified or isolated polypeptide encoded by a polynucleotide of the present invention. In a non-limiting embodiment, the polypeptide is a HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 *L. intracellularis* protein. In a preferred embodiment, the *L. intracellularis* HtrA, PonA, HypC, LysS, YcfW, ABC1, and Omp100 proteins have the amino acid sequences of SEQ ID NOS: 3-8 or SEQ ID NO: 102. In another embodiment, the polypeptides are substantially free of other *Lawsonia* proteins.

[0117] The present invention further provides polypeptides that are homologous to any of the aforementioned L. intracellularis proteins, as the term "homologous" is defined above for polypeptides. Polypeptides of the present invention that are homologous to the proteins of the invention do not include polypeptides having the amino acid sequences of non-Lawsonia proteins described herein. The polypeptide of the invention, in one embodiment, has more than 70%, preferably more than about 90%, and most preferably more than about 95% amino acid sequence identity to the Lawsonia proteins, wherein sequence identity is determined by use of the BLASTP algorithm (GenBank, NCBI).

[0118] In another embodiment, the polypeptide consists of the *L. intracellularis* HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein having between 1 and 10, and more preferably between 1 and 5, amino acids inserted, deleted, or substituted, including combinations thereof. In a more particular example of this embodiment, the isolated polypeptide has between 1 and 5 amino acids conservatively substituted in the amino acid sequence of the HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein.

[0119] The present invention further provides polypeptides consisting of a substantial portion of any one of the aforementioned polypeptides of the present invention. As used herein, a "substantial portion" of a polypeptide of the present invention, or "peptide fragment," means a polypeptide consisting of less than the complete amino acid sequence of the corresponding full-length polypeptide, but comprising at least about 5%, more preferably at least about 20%, even more preferably at least about 50%, and most preferably at least about 95% of the amino acid sequence thereof, and that is useful in practicing the present invention. Particularly preferred are peptide fragments that are immunogenic, i.e., capable of inducing an immune response which results in production of antibodies that react specifically against the corresponding full-length Lawsonia polypeptide.

[0120] In another embodiment, the polypeptide of the invention comprises an epitope of HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein that is specifically reactive with anti-Lawsonia antibodies. The epitope is preferably more than 8, more preferably more than 12, and most preferably, more than 20 amino ocids of the protein sequence.

[0121] The present invention further provides fusion proteins comprising any of the polypeptides of the invention fused to a carrier or fusion partner as known in the art.

[0122] The present invention further provides a method of preparing any of the polypeptides described above, comprising culturing a host cell transformed with a recombinant expression vector, said recombinant expression vector comprising a polynucleotide molecule comprising a nucleotide sequence encoding the particular polypeptide, which polynucleotide molecule is in operative association with one or more regulatory elements, under conditions conducive to the expression of the polypeptide, and recovering the expressed polypeptide from the cell culture.

Use Of Polypeptides

Once a polypeptide of the present invention of sufficient purity has been obtained, it can be characterized by standard methods, including by SDS-PAGE, size exclusion chromatography, amino acid sequence analysis, immunological activity, biological activity, etc. The polypeptide can be further characterized using hydrophilicity analysis (see, e.g., Hopp and Woods, 1981, Proc. Natl. Acad. Sci. USA 78:3824), or analogous software algorithms, to identify hydro-

phobic and hydrophilic regions. Structural analysis can be carried out to identify regions of the polypeptide that assume specific secondary structures. Biophysical methods such as X-ray crystallography (Engstrom, 1974, Biochem. Exp. Biol. 11: 7-13), computer modeling (Fletterick and Zoller (eds), 1986, in: <u>Current Communications in Molecular Biology</u>. Cold Spring Harbor, Cold Spring Harbor, NY), and nuclear magnetic resonance (NMR) can be used to map and study potential sites of interaction between the polypeptide and other putative interacting proteins/receptors/molecules. Information obtained from these studies can be used to design deletion mutants and vaccine compositions, and polypeptide in vivo.

[0124] Polypeptides of the present invention are useful for a variety of purposes, including as components of vaccine compositions to protect PPE susceptible animals against PPE; or as diagnostic reagents, e.g., using standard techniques such as ELISA assays, to screen for *Lawsonia*-specific antibodies in blood or serum samples from animals; or as antigens to raise polyclonal or monoclonal antibodies, as described below, which antibodies are useful as diagnostic reagents, e.g., using standard techniques such as Western blot assays, to screen for *Lawsonia*-specific proteins in cell, tissue or fluid samples from an animal.

Analogs And Derivatives Of Polypeptides

[0125] A polypeptide of the present invention can be modified at the protein level to improve or otherwise alter its biological or immunological characteristics. One or more chemical modifications of the polypeptide can be carried out using known techniques to prepare analogs therefrom, including but not limited to any of the following: substitution of one or more L-amino acids of the polypeptide with corresponding D-amino acids, amino acid analogs, or amino acid mimics, so as to produce, e.g., carbazates or tertiary centers; or specific chemical modification, such as, e.g., proteo-acetylation, formylation, oxidation or reduction, etc. Alternatively or additionally, polypeptides of the present invention [0126]

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[0126] A polypeptide of the present invention can be derivatized by conjugation thereto of one or more chemical groups, including but not limited to acetyl groups, sulfur bridging groups, glycosyl groups, lipids, and phosphates, and/or by conjugation to a second polypeptide of the present invention, or to another protein, such as, e.g., serum albumin, keyhole limpet hemocyanin, or commercially activated BSA, or to a polyamino acid (e.g., polylysine), or to a polysacharide, (e.g., sepharose, agarose, or modified or unmodified celluloses), among others. Such conjugation is preferably by covalent linkage at amino acid side chains and/or at the N-terminus or C-terminus of the polypeptide. Methods for carrying out such conjugation reactions are well-known in the field of protein chemistry.

[0127] Derivatives useful in practicing the claimed invention also include those in which a water-soluble polymer such as, *e.g.*, polyethylene glycol, is conjugated to a polypeptide of the present invention, or to an analog or derivative thereof, thereby providing additional desirable properties while retaining, at least in part, the immunogenicity of the polypeptide. These additional desirable properties include, *e.g.*, increased solubility in aqueous solutions, increased stability in storage, increased resistance to proteolytic dehydration, and increased *in vivo* half-life. Water-soluble polynomopolymers suitable for conjugation to a polypeptide of the present invention include but are not limited to polyethylene glycol homopolymers, polypropylene glycol homopolymers of ethylene glycol with propylene glycol, wherein said homopolymers and copolymers are unsubstituted or substituted at one end with an alkyl group, polyoxyethylated polyols, polyvinyl alcohol, polysaccharides, polyvinyl ethyl ethers, and α,β-poly[2-hydroxyethyl]-DL-aspartamide. Polyethylene glycol is particularly preferred. Methods for making water-soluble polymer conjugates of polypeptides are known in the art and are described in, among other places, U.S. Patent 3,788,948; U.S. Patent 3,960,830; U.S. Patent 4,002,531; U.S. Patent 4,055,635; U.S. Patent 4,179,337; U.S. Patent 4,261,973; U.S. Patent 4,412,989; U.S. Patent 4,414,147; U.S. Patent 4,415,665; U.S. Patent 4,609,546; U.S. Patent 4,732,863; U.S. Patent 4,745,180; European Patent (EP) 152,847; EP 98,110; and Japanese Patent 5,792,435, which patents are incorporated herein by reference.

Antibodies

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50 [0128] The present invention further provides isolated antibodies directed against a polypeptide of the present invention. In a preferred embodiment, antibodies can be raised against a HtrA, PonA, HypC, LysS, YcfW, ABC1, or rabbits, goats, sheep, or mice, can be immunized with a partially or substantially purified, or isolated, *L. intracellularis* protein, or with a homolog, fusion protein, substantial portion, analog or derivative thereof, as these are described above. An adjuvant, such as described below, can be used to enhance antibody production.

[0129] Polyclonal antibodies can be obtained and isolated from the serum of an immunized animal and tested for specificity against the antigen using standard techniques. Alternatively, monoclonal antibodies can be prepared and isolated using any technique that provides for the production of antibody molecules by continuous cell lines in culture.

These include but are not limited to the hybridoma technique originally described by Kohler and Milstein (Nature, 1975, 256: 495-497); the human B-cell hybridoma technique (Kosbor *et al.*, 1983, Immunology Today 4:72; Cote *et al.*, 1983, Proc. Natl. Acad. Sci. *USA* 80: 2026-2030); and the EBV-hybridoma technique (Cole *et al.*, 1985, Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, Inc., pp. 77-96). Alternatively, techniques described for the production of single chain antibodies (see, *e.g.*, U.S. Patent 4,946,778) can be adapted to produce *L. intracellularis* antigen-specific single chain antibodies. These publications are incorporated herein by reference.

[0130] Antibody fragments that contain specific binding sites for a polypeptide of the present invention are also encompassed within the present invention, and can be generated by known techniques. Such fragments include but are not limited to F(ab')₂ fragments, which can be generated by pepsin digestion of an intact antibody molecule, and Fab fragments, which can be generated by reducing the disulfide bridges of the F(ab')₂ fragments. Alternatively, Fab expression libraries can be constructed (Huse *et al.*, 1989, Science 246: 1275-1281) to allow rapid identification of Fab fragments having the desired specificity to the *L. intracellularis* protein.

[0131] Techniques for the production and isolation of monoclonal antibodies and antibody fragments are well-known in the art, and are additionally described, among other places, in Harlow and Lane, 1988, Antibodies: A Laboratory Manual, Cold Spring Harbor Laboratory, and in J. W. Goding, 1986, Monoclonal Antibodies: Principles and Practice, Academic Press, London, which are incorporated herein by reference.

Targeted Mutation Of Lawsonia Genes

[0132] Based on the disclosure of the polynucleotide molecules of the present invention, genetic constructs can be prepared for use in disabling or otherwise mutating a Lawsonia htrA, ponA, lysS, ycfW, hypC, abc1, or omp100 gene (which gene is hereinafter referred to as the "Lawsonia gene"). The Lawsonia gene can be mutated using an appropriately designed genetic construct. For example, the Lawsonia gene can be mutated using a genetic construct of the present invention that functions to: (a) delete all or a portion of the coding sequence or regulatory sequence of the Lawsonia gene with a different nucleotide sequence; or (c) insert into the coding sequence or regulatory sequence of the Lawsonia gene one or more nucleotides, or an oligonucleotide molecule, or polynucleotide molecule, which can comprise a nucleotide sequence from Lawsonia or from a heterologous source; or (d) carry out some combination of (a), (b) and (c). Alternately, constructs can be employed to alter the expression of the Lawsonia gene or the stability of its encoded protein.

[0133] Lawsonia cells in which a Lawsonia gene has been mutated are, for example, useful in practicing the present invention where mutating the gene reduces the pathogenicity of the Lawsonia cells carrying the mutated gene compared to cells of the same strain of Lawsonia where the gene has not been so mutated, and where such Lawsonia cells carrying the disabled gene can be used in a vaccine composition, particularly in a modified live vaccine, to induce or contribute to the induction of, a protective response in an animal against PPE. In a preferred embodiment, the mutation serves to partially or completely disable the Lawsonia gene, or partially or completely disable the protein encoded by the Lawsonia gene. In this context, a Lawsonia gene or protein is considered to be partially or completely disabled if either no protein product is made (for example, the gene is deleted), or a protein product is made that can no longer carry out its normal biological function or can no longer be transported to its normal cellular location, or a product is made that carries out its normal biological function but at a significantly reduced rate. Lawsonia cells in which the Lawsonia gene has been mutated are also useful to increase expression of that gene or the stability of its encoded protein. Mutations are particularly useful that result in a detectable decrease in the pathogenicity of cells of a pathogenic strain of Lawsonia. The invention also encompasses cells expressing proteins and polypeptides of the invention where such cells are constitutive mutants.

[0134] In a non-limiting embodiment, a genetic construct of the present invention is used to mutate a wild-type Lawsonia gene by replacement of the coding sequence of the wild-type gene, or a promoter or other regulatory region thereof, or a portion thereof, with a different nucleotide sequence such as, e.g., a mutated coding sequence or mutated regulatory region, or portion thereof. Mutated Lawsonia gene sequences for use in such a genetic construct can be produced by any of a variety of known methods, including by use of error-prone PCR, or by cassette mutagenesis. For example, oligonucleotide-directed mutagenesis can be employed to after the coding sequence or promoter sequence of a wild-type Lawsonia gene in a defined way, e.g., to introduce a frame-shift or a termination codon at a specific point within the sequence. Alternatively or additionally, a mutated nucleotide sequence for use in the genetic construct of the present invention can be prepared by insertion or deletion of the coding sequence or promoter sequence of one or more nucleotides, oligonucleotide molecules or polynucleotide molecules, or by replacement of a portion of the coding sequence or promoter sequence with one or more different nucleotides, oligonucleotide molecules or polynucleotide molecules can be obtained from any naturally occurring source or can be synthetic. The inserted or deleted sequence can serve simply to disrupt the reading frame of the Lawsonia gene, or can further encode a heterologous gene product such as a selectable marker.

[0135] Alternatively or additionally, random mutagenesis can be used to produce a mutated *Lawsonia* gene sequence for use in a genetic construct of the present invention. Random mutagenesis can be carried out by any suitable techniques such as, *e.g.*, by exposing cells carrying a *Lawsonia* gene to ultraviolet radiation or x-rays, or to chemical mutagens such as N-methyl-N'-nitrosoguanidine, ethyl methane sulfonate, nitrous acid or nitrogen mustards, and then selecting for cells carrying a mutation in the particular gene. See, *e.g.*, Ausubel, 1989, above, for a review of mutagenesis techniques.

[0136] Mutations to produce modified *Lawsonia* cells that are useful in practicing the present invention can occur anywhere in the *Lawsonia* gene, including in the ORF, or in the promoter or other regulatory region, or in any other sequences that naturally comprise the gene or ORF, or that alter expression of the gene or the stability of its encoded protein. Such *Lawsonia* cells include mutants in which a modified form of the protein normally encoded by the *Lawsonia* gene is produced, or in which no protein normally encoded by the *Lawsonia* gene is produced, and can be null, conditional, constitutive, or leaky mutants.

[0137] Alternatively, a genetic construct of the present invention can comprise nucleotide sequences that naturally flank the *Lawsonia* gene or ORF *in situ*, with only a portion or no nucleotide sequences from the coding region of the gene itself. Such a genetic construct would be useful, *e.g.*, to delete the entire *Lawsonia* gene or ORF.

[0138] In one embodiment, a genetic construct of the present invention comprises a polynucleotide molecule that can be used to disable a *Lawsonia* gene, comprising: (a) a polynucleotide molecule having a nucleotide sequence that is otherwise the same as a nucleotide sequence encoding a HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein from *L. intracellularis*, but which nucleotide sequence further comprises one or more disabling mutations; or (b) a polynucleotide molecule comprising a nucleotide sequence that naturally flanks the ORF of a *Lawsonia* gene *in situ*. Once transformed into cells of a strain of *Lawsonia*, the polynucleotide molecule of the genetic construct is specifically targeted to the particular *Lawsonia* gene by homologous recombination, and thereby either replaces the gene or portion thereof or inserts into the gene. As a result of this recombination event, the *Lawsonia* gene otherwise native to that particular strain of *Lawsonia* is disabled.

[0139] In another embodiment, a genetic construct employs a mutation that alters expression, e.g., by constitutively expressing or overexpressing the HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein. Such a mutation can be useful, for example, to weaken the host cells. The construct can also employ a mutation that increases stability of the protein to, e.g., attenuate the host cell.

[0140] For targeted gene mutation through homologous recombination, the genetic construct is preferably a plasmid, either circular or linearized, comprising a mutated nucleotide sequence as described above. In a non-limiting embodiment, at least about 200 nucleotides of the mutated sequence are used to specifically direct the genetic construct of the present invention to the particular targeted *Lawsonia* gene for homologous recombination, although shorter lengths of nucleotides can also be effective. In addition, the plasmid preferably comprises an additional nucleotide sequence encoding a reporter gene product or other selectable marker that is constructed so that it will insert into the *Lawsonia* genome in operative association with the regulatory element sequences of the native *Lawsonia* gene to be disrupted. Reporter genes that can be used in practicing the invention are well-known in the art and include those encoding CAT, green fluorescent protein, and β-galactosidase, among others. Nucleotide sequences encoding selectable markers are also well-known in the art, and include those that encode gene products conferring resistance to antibiotics or anti-metabolites, or that supply an auxotrophic requirement. Examples of such sequences include those that encode pyrimethamine resistance, or neomycin phosphotransferase (which confers resistance to hygromycin).

[0141] Methods that can be used for creating the genetic constructs of the present invention are well-known in the art, and include *in vitro* recombinant techniques, synthetic techniques, and *in vivo* genetic recombination, as described, among other places, in Maniatis *et al.*, 1989, above; Ausubel *et al.*, 1989, above; Sambrook *et al.*, 1989, above; Innis *et al.*, 1995, above; and Erlich, 1992, above.

[0142] Lawsonia cells can be transformed or transfected with a genetic construct of the present invention in accordance with known techniques, such as, e.g., by electroporation. Selection of transformants can be carried out using standard techniques, such as by selecting for cells expressing a selectable marker associated with the construct. Identification of transformants in which a successful recombination event has occurred and the particular target gene has been altered can be carried out by genetic analysis, such as by Southern blot analysis, or by Northern analysis to detect a lack of mRNA transcripts encoding the particular protein, or cells lacking the particular protein, as determined, e.g., by immunological analysis, by the appearance of a novel phenotype, such as reduced pathogenicity, by PCR assay, or by some combination thereof.

[0143] In a further non-limiting embodiment, the genetic construct of the present invention can additionally comprise a different gene or coding region from Lawsonia or from a different pathogen that infects the animal, which gene or coding region encodes an antigen useful to induce, or contribute to the induction of, a separate and distinct protective immune response in the animal upon vaccination with the modified live Lawsonia cells of the present invention. This additional gene or coding region can be further engineered to contain a signal sequence that leads to secretion of the

encoded antigen from the modified live *Lawsonia* cell, thereby allowing for the antigen to be displayed to the immune system of the vaccinated animal.

[0144] The present invention thus provides modified live Lawsonia cells in which the htrA, ponA, hypC, lysS, ycfW, abc1, or omp100 gene has been mutated. In addition, the present invention provides a method of preparing modified live Lawsonia cells, comprising: (a) transforming cells of Lawsonia with a genetic construct of the invention; (b) selecting transformed cells in which the htrA, ponA, hypC, lysS, ycfW abc1, or omp100 gene has been mutated by the genetic construct; and (c) selecting from among the cells of step (b) those cells that can be used in a vaccine to protect a PPE susceptible animal against PPE. The invention also encompasses killed cell compositions prepared from such modified Lawsonia cells.

Culturing Lawsonia Bacteria

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[0145] Lawsonia bacterium for use in the present invention can be cultured and maintained in vitro using methods described e.g. by Joens et al., 1997, Am. J. Vet. Res. 58:1125-1131; Lawson et al., 1993, Journal of Clinical Microbiology 31:1136-1142; and McOrist et al., 1995, supra.

Anti-Lawsonia Vaccines

[0146] The present invention further provides a vaccine against PPE, comprising an immunologically effective amount of a protein or polypeptide of the present invention, and a pharmaceutically acceptable carrier. In a preferred embodiment, the vaccine comprises a HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 *L. intracellularis* protein.

[0147] The present invention further provides a vaccine against PPE, comprising an immunologically effective amount of one or more polynucleotide molecules of the present invention, and a pharmaceutically acceptable carrier. In a preferred embodiment, the vaccine comprises a polynucleotide molecule having a nucleotide sequence encoding *L. intracellularis* HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100.

[0148] The present invention further provides a vaccine against PPE, comprising an immunologically effective amount of modified *Lawsonia* bacteria of the present invention, and a pharmaceutically acceptable carrier. In one embodiment, the modified *Lawsonia* cells for use in the vaccine of the present invention are live *L. intracellularis* bacteria which express a HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 phenotype. Alternatively, the vaccine of the present invention can comprise any of such modified *Lawsonia* cells of the present invention that have been inactivated. Inactivation of modified *Lawsonia* cells can be carried out using any techniques known in the art, including by chemical treatment, such as with binary ethylenimine (BEI), or beta-propiolactone, or formaldehyde, or by freeze-thawing or heat treatment, or by homogenization of cells, or by a combination of these types of techniques. Vaccines prepared from homogenized, modified *Lawsonia* cells can consist of either the entire unfractionated cell homogenate, or an immunologically effective subfraction thereof.

[0149] As used herein, the term "immunologically effective amount" refers to that amount of antigen, e.g., protein, polypeptide, polynucleotide molecule, or modified cells, capable of inducing a protective response against PPE when administered to a member of a PPE susceptible animal species after either a single administration, or after multiple administrations.

[0150] The phrase "capable of inducing a protective response" is used broadly herein to include the induction or enhancement of any immune-based response in the animal in response to vaccination, including either an antibody or cell-mediated immune response, or both, that serves to protect the vaccinated animal against PPE. The terms "protective response" and "protect" as used herein to refer not only to the absolute prevention of PPE or absolute prevention of infection by Lawsonia, but also to any detectable reduction in the degree or rate of infection by such a pathogen, or any detectable reduction in the severity of the disease or any symptom or condition resulting from infection by the pathogen, including, e.g., any detectable reduction in the rate of formation, or in the absolute number, of lesions formed in one or more tissues, or the transmission of infection to other animals, in the vaccinated animal as compared to an unvaccinated infected animal of the same species.

[0151] In a further preferred embodiment, the vaccine of the present invention is a combination vaccine for protecting a PPE susceptible animal against PPE and, optionally, one or more other diseases or pathological conditions that can afflict the animal, which combination vaccine comprises an immunologically effective amount of a first component comprising a polypeptide, polynucleotide molecule, or modified Lawsonia cells of the present invention; an immunologically effective amount of a second component that is different from the first component, and that is capable of inducing, or contributing to the induction of, a protective response against a disease or pathological condition that can afflict the PPE susceptible animal; and a pharmaceutically acceptable carrier.

[0152] The second component of the combination vaccine is selected based on its ability to induce, or contribute to the induction of, a protective response against either PPE or another disease or pathological condition that can afflict members of the relevant species, as known in the art. Any antigenic component that is useful in a vaccine composition

in the particular species can be used as the second component of the combination vaccine. Such antigenic components include but are not limited to those that provide protection against pathogens selected from the group consisting of Leptospira spp., Campylobacter spp., Staphylococcus aureus, Streptococcus agalactiae, Streptococcus suis, Mycoplasma spp., Klebsiella spp., Salmonella spp., rotavirus, coronavirus, rabies, Pasteurella hemolytica, Pasteurella multocida, Clostridia spp., Tetanus toxoid, E. coli, Cryptosporidium spp., Eimeria spp., Trichomonas spp., Serpulina (Brachyspira) hyodysenteriae, Actinobacillus pleuropneumoniae, Actinobacillus suis, Salmonella cholerasuis, Erysipelothrix rhusiopathiae, Leptospira sp., Staphylococcus hyicus, Haemophilus parasuis, Bordetella bronchiseptica, Mycoplasma hyopneumoniae, porcine reproductive and respiratory syndrome virus, swine influence virus, porcine immunodeficiency virus, transmissible gastroenteritis virus, porcine parvovirus, encophalomyocarditis virus, coronavirus, pseudorabies virus, circovirus and other eukarvotic parasites.

[0153] In one embodiment, the combination vaccine of the present invention comprises a combination of two or more components selected from the group consisting of an immunologically effective amount of a protein or polypeptide of the present invention, an immunologically effective amount of a polynucleotide molecule of the present invention, and an immunologically effective amount of modified *Lawsonia* cells of the present invention.

[0154] The vaccines of the present invention can further comprise one or more additional immunomodulatory components including, e.g., an adjuvant or cytokine, as described below.

[0155] The present invention further provides a method of preparing a vaccine against PPE, comprising combining an immunologically effective amount of a *L. intracellularis* protein or polypeptide, or polynucleotide molecule, or modified *Lawsonia* cells, of the present invention, with a pharmaceutically acceptable carrier, in a form suitable for administration to a PPE susceptible animal. In a preferred embodiment, the protein is *L. intracellularis* HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100, the polynucleotide molecule preferably comprises a nucleotide sequence encoding *L. intracellularis* HtrA, PonA, HypC, LysS. YcfW, ABC1, or Omp100 protein and the modified *Lawsonia* bacteria has an HtrA', PonA', HypC', LysS', YcfW', ABC1', or Omp100' phenotype.

[0156] A vaccine comprising modified live *Lawsonia* cells of the present invention can be prepared using an aliquot of culture fluid containing said *Lawsonia* cells, either free in the medium or residing in mammalian host cells, or both, and can be administered directly or in concentrated form to the PPE susceptible animal. Alternatively, modified live *Lawsonia* cells can be combined with a pharmaceutically acceptable carrier, with or without an immunomodulatory agent, selected from those known in the art and appropriate to the chosen route of administration, preferably where at least some degree of viability of the modified live *Lawsonia* cells in the vaccine composition is maintained.

Vaccine compositions of the present invention can be formulated following accepted convention to include pharmaceutically acceptable carriers, such as standard buffers, stabilizers, diluents, preservatives, and/or solubilizers, and can also be formulated to facilitate sustained release. Diluents include water, saline, dextrose, ethanol, glycerol, and the like. Additives for isotonicity include sodium chloride, dextrose, mannitol, sorbitol, and lactose, among others. Stabilizers include albumin, among others. Suitable other vaccine vehicles and additives, including those that are particularly useful in formulating modified him vaccines are known as will be appropriate the second additives.

ticularly useful in formulating modified live vaccines, are known or will be apparent to those skilled in the art,. See, e.g., Remington's Pharmaceutical Science, 18th ed., 1990, Mack Publishing, which is incorporated herein by reference.

[0158] The vaccine of the present invention can further comprise one or more additional immunomodulatory com-

ponents such as, *e.g.*, an adjuvant or cytokine, among others. Non-limiting examples of adjuvants that can be used in the vaccine of the present invention include the RIBI adjuvant system (Ribi Inc., Hamilton, MT), alum, mineral gels such as aluminum hydroxide gel, oil-in-water emulsions, water-in-oil emulsions such as, *e.g.*, Freund's complete and incomplete adjuvants, Block co polymer (CytRx, Atlanta GA), QS-21 (Cambridge Biotech Inc., Cambridge MA), SAF-M (Chiron, Emeryville CA), AMPHIGEN[®] adjuvant, saponin, Quil A or other saponin fraction, monophosphoryl lipid A, Avridine lipid-amine adjuvant, and protein adjuvants such as *Vibrio cholera* toxin and *E. coli* labile toxin. Specific non-limiting examples of oil-in-water emulsions useful in the vaccine of the invention include modified SEAM62 and SEAM 1/2 formulations. Modified SEAM62 is an oil-in-water emulsion containing 5% (v/v) squalene (Sigma), 1% (v/v) SPAN[®] 85 detergent (ICI Surfactants), 2.5% (v/v) ethanol, 200 μg/ml Quil A, 100 μg/ml cholesterol, and 0.5% (v/v) lecithin. Modified SEAM 1/2 is an oil-in-water emulsion comprising 5% (v/v) squalene, 1% (v/v) SPAN[®] 85 detergent, 0.7% (v/v) Tween 80 detergent, 2.5% (v/v) ethanol, 100 μg/ml Quil A, and 50 μg/ml cholesterol. Other immunomodulatory agents that can be included in the vaccine include, e.g., one or more interleukins, interferons, or other known cytokines. Where the vaccine comprises modified live *Lawsonia* cells, the adjuvant is preferably selected based on the ability of the resulting vaccine formulation to maintain at least some degree of viability of the modified live *Lawsonia* cells.

[0159] Where the vaccine composition comprises a polynucleotide molecule, the polynucleotide molecule can either be DNA or RNA, although DNA is preferred, and is preferably administered to a PPE susceptible animal to be protected against PPE in an expression vector construct, such as a recombinant plasmid or viral vector, as known in the art. Examples of recombinant viral vectors include recombinant adenovirus vectors and recombinant retrovirus vectors. The vaccine formulation can also comprise a non-viral DNA vector, such as a DNA plasmid-based vector. The polynucleotide molecule may be associated with lipids to form, e.g., DNA-lipid complexes, such as liposomes or coch-

leates. See, e.g., International Patent Publication WO 93/24640.

[0160] An expression vector useful as a vaccinal agent in a DNA vaccine preferably comprises a nucleotide sequence encoding one or more antigenic *Lawsonia* proteins, or a substantial portion of such a nucleotide sequence, in operative association with one or more transcriptional regulatory elements required for expression of the *Lawsonia* coding sequence in a eukaryotic cell, such as, *e.g.*, a promoter sequence, as known in the art. In a preferred embodiment, the regulatory element is a strong viral promoter such as, *e.g.*, a viral promoter from RSV or CMV. Such an expression vector also preferably includes a bacterial origin of replication and a prokaryotic selectable marker gene for cloning purposes, and a polyadenylation sequence to ensure appropriate termination of the expressed mRNA. A signal sequence may also be included to direct cellular secretion of the expressed protein.

[0161] The requirements for expression vectors useful as vaccinal agents in DNA vaccines are further described in U.S. Patent 5,703,055, U.S. Patent 5,580,859, U.S. Patent 5,589,466, International Patent Publication WO 98/35562, and in various scientific publications, including Ramsay *et al.*, 1997, Immunol. Cell Biol. 75:360-363; Davis, 1997, Cur. Opinion Biotech. 8:635-640: Maniackan *et al.*, 1997, Critical Rev. Immunol. 17:139-154; Robinson, 1997, Vaccine 15(8):785-787; Lai and Bennett, 1998, Critical Rev. Immunol. 18:449-484; and Vogel and Sarver, 1995, Clin. Microbiol. Rev. 8(3):406-410, among others.

[0162] Where the vaccine composition comprises modified live *Lawsonia* cells, the vaccine can be stored cold, frozen, or lyophilized. Where the vaccine composition instead comprises a protein, polypeptide, polynucleotide molecule, or inactivated modified *Lawsonia* cells of the present invention, the vaccine may be stored cold, frozen, or in lyophilized form to be rehydrated prior to administration using an appropriate diluent.

[0163] The vaccine of the present invention can optionally be formulated for sustained release of the antigen. Examples of such sustained release formulations include antigen in combination with composites of biocompatible polymers, such as, e.g., poly(lactic acid), poly(lactic-co-glycolic acid), methylcellulose, hyaluronic acid, collagen and the like. The structure, selection and use of degradable polymers in drug delivery vehicles have been reviewed in several publications, including A. Domb et al., 1992, Polymers for Advanced Technologies 3: 279-292, which is incorporated herein by reference. Additional guidance in selecting and using polymers in pharmaceutical formulations can be found in the text by M. Chasin and R. Langer (eds), 1990, "Biodegradable Polymers as Drug Delivery Systems" in: <u>Drugs and the Pharmaceutical Sciences</u>, Vol. 45, M. Dekker, NY, which is also incorporated herein by reference. Alternatively, or additionally, the antigen can be microencapsulated to improve administration and efficacy. Methods for microencapsulating antigens are well-known in the art, and include techniques described, e.g., in U.S. Patent 3,137,631; U.S. Patent 3,959,457; U.S. Patent 4,205,060; U.S. Patent 4,606,940; U.S. Patent 4,744,933; U.S. Patent 5,132,117; and International Patent Publication WO 95/28227, all of which are incorporated herein by reference.

[0164] Liposomes can also be used to provide for the sustained release of antigen. Details concerning how to make and use liposomal formulations can be found in, among other places, U.S. Patent 4,016,100; U.S. Patent 4,452,747; U.S. Patent 4,921,706; U.S. Patent 4,927,637; U.S. Patent 4,944,948; U.S. Patent 5,008,050; and U.S. Patent 5,009,956, all of which are incorporated herein by reference.

[0165] The present invention further provides a method of vaccinating a PPE susceptible animal against PPE, comprising administering to the animal an immunogenically effective amount of a vaccine of the present invention. The vaccine is preferably administered parenterally, e.g., either by subcutaneous or intramuscular injection. However, the vaccine can also be administered by intraperitoneal or intravenous injection, or by other routes, including, e.g., orally, intranasally, rectally, vaginally, intra-ocularly, or by a combination of routes, and also by delayed release devices as known in the art. The skilled artisan can determine optimal routes of vaccine administration, and recognize acceptable formulations for the vaccine composition according to the chosen route of administration.

[0166] An effective dosage can be determined by conventional means, starting with a low dose of antigen, and then increasing the dosage while monitoring its effects. Numerous factors may be taken into consideration when determining an optimal dose per animal. Primary among these is the species, size, age and general condition of the animal, the presence of other drugs in the animal, the virulence of a particular strain of *Lawsonia* against which the animal is being vaccinated, and the like. The actual dosage is preferably chosen after consideration of the results from other animal studies.

[0167] The dose amount of a protein or polypeptide of the present invention in a vaccine of the present invention preferably ranges from about 1 μg to about 10 mg, more preferably from about 50 μg to about 1 mg, and most preferably from about 100 μg to about 0.5 mg. The dose amount of a *Lawsonia* polynucleotide molecule of the present invention in a vaccine of the present invention preferably ranges from about 50 μg to about 1 mg. The dose amount of modified *Lawsonia* cells of the present invention in a vaccine of the present invention preferably ranges from about 1 x 10³ to about 1 x 10⁸ cells/ml, and more preferably from about 1 x 10⁵ to about 1 x 10⁷ cells/ml. A suitable dosage size ranges from about 0.1 ml to about 10 ml, and more preferably from about 1 ml to about 5 ml. The dose amounts of these antigens are also applicable to combination vaccines of the present invention. Where the second component of the combination vaccine is an antigen other than a *Lawsonia* protein, polypeptide, polynucleotide or modified cell of the present invention, the dose amount of the second component for use in the combination vaccine can be determined from prior

vaccine applications of that second component, as known in the art.

[0168] The vaccine of the present invention is useful to protect animals, especially pigs, against PPE. The vaccine can be administered to any suitable animals, including, without limitation, hamsters, ferrets, guinea pigs, deer, and bovine, equine, and avian species. The vaccine of the invention can be administered at any time during the life of a particular animal depending upon several factors including, e.g., the timing of an outbreak of PPE among other animals, etc. The vaccine can be administered to animals of weaning age or younger, or to more mature animals. Effective protection may require only a primary vaccination, or one or more booster vaccinations may also be needed. One method of detecting whether adequate immune protection has been achieved is to determine seroconversion and antibody titer in the animal after vaccination. The timing of vaccination and the number of boosters, if any, is preferably determined by a veterinarian based on analysis of all relevant factors, some of which are described above.

[0169] In one embodiment, a protein or polypeptide of the invention, e.g., HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 *L. intracellularis* protein, or combinations thereof, is administered in a formulation containing 100µg of polypeptide, and 25µg of *E. coli* labile toxin as adjuvant, in 1 ml of buffered solution. The formulation is, for example, administered intramuscularly to pigs at between 5 and 7 days of age, and readministered 14 days later.

[0170] The present invention further provides a kit for vaccinating a PPE susceptible animal against PPE, comprising a container having an immunologically effective amount of a polypeptide, polynucleotide molecule, or modified Lawsonia cells of the present invention, or a combination thereof. The kit can optionally comprise a second container having a pharmaceutically acceptable carrier or diluent. In a preferred embodiment, the polypeptide is the HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 L. intracellularis protein; the polynucleotide molecule preferably has a nucleotide sequence that encodes the HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 L. intracellularis protein and the modified Lawsonia cells preferably are live or inactivated cells that express an HtrA⁻, PonA⁻, HypC⁻, LysS⁻, YcfW, ABC1⁻, or Omp100 phenotype.

[0171] The invention also relates to a kit for detecting the presence of *L. intracellularis*, an *L. intracellularis* specific amino acid or nucleotide sequence, or an anti- *L. intracellularis* antibody, that contains a protein, polypeptide, polynucleotide, or antibody of the invention. The kit can also contain means for detecting the protein, polypeptide, polynucleotide, or antibody of the invention including, for example, an enzyme, fluorescent, or radioactive label attached to the protein, polypeptide, polynucleotide, or antibody, or attached to a moiety that binds to the protein, polypeptide, polynucleotide, or antibody.

[0172] The following examples are illustrative only, and not intended to limit the scope of the present invention.

EXAMPLES

Example 1: Molecular cloning of L. intracellularis chromosomal Region A

35 Isolation of DNA and construction of DNA libraries

[0173] Template DNA was purified from pig intestinal mucosa isolated from the ileum of pigs experimentally infected with *L. Intracellularis*. DNA purification from homogenized Intestinal mucosa was performed according to (1) the method of Nollau *et al.* (Nollau et al., 1996, BioTechniques 20: 784-788) or (2) phenol extraction and sodium acetate-ethanol precipitation of DNA. To facilitate cloning of *L. intracellularis* gene sequences, several genomic libraries were constructed. These libraries were specifically modified by ligation of a known sequence (Vectorette II™, Genosys Biotechnologies, Inc., The Woodlands, TX) to the 5' and 3' ends of restricted DNA fragments. Vectorette™ libraries were constructed by separately digesting aliquots of *L. intracellularis*-infected pig mucosal DNA extract with restriction endonucleases *Hind*III, *Eco*RI, *Dra*I or *Hpa*I at 37° overnight. The reaction was then spiked with additional fresh restriction enzyme and adjusted to 2 mM ATP, 2 mM DTT final concentration. Vectorette™ tailing was carried out by addition of T₄ DNA Ligase (400 U) plus 3 pMoI of the appropriate compatible Vectorette™ linker (*Hind*III Vectorette™: *Hind*III digested DNA; *Eco*RI: *Eco*RI digested DNA; Blunt: *Dra*I, *Hpa*I digested DNA). The mixture was incubated for three cycles at 20°, 60 min; and 37°, 30 min to complete the tailing reaction then adjusted to 200 μl with water and stored at -20°.

50 Molecular cloning of genomic Region A encoding LysS, YcfW, ABC1, and Omp100 proteins

[0174] Identification of genomic Region A (shown in Fig. 1) was accomplished by genomic walking and phage library screening processes. Screening of the *HindIII*, *EcoRI*, *DraI*, and *HpaI* Vectorette™ libraries was carried out to obtain DNA fragments located adjacent to gene (*amiB*) from *L. intracellularis* having homology to bacterial N-acetyl-muramoyl-L-alanine amidases involved in cell wall autolysis. Oligonucleotide primers ER159 (SEQ ID NO: 37), ER161 (SEQ ID NO: 38), and ER162 (SEQ ID NO: 39) were designed based on the nucleotide sequence of *amiB*. All three primers were designed to bind the (-) strand within this region to allow amplification of DNA located upstream of the gene encoding AmiB.

[0175] For polymerase chain amplification of a fragment of the *omp100* gene, oligonucleotides ER159 (SEQ ID NO: 37), ER161 (SEQ ID NO: 38), and ER162 (SEQ ID NO: 39) were used in combination with a Vectorette[™] specific primer (ER70) (SEQ ID NO: 33) in 50 μl reactions containing 1x PCR Buffer II (Perkin Elmer), 1.5 mM MgCl₂, 200 μM each deoxy-NTP, 50 pMol each primer, and 2.5 U AmpliTaq[™] Gold (Perkin Elmer) thermostable polymerase. Multiple single reactions were performed with 4 μl of the Vectorette[™] libraries as DNA template. Amplification was carried out as follows: denaturation (95° 9 min); 40 cycles of denaturation (95° 30 sec), annealing (65° 30 sec), and polymerization (72° 2.5 min); followed by a final extension at 72° for 7 minutes.

[0176] The amplified products were visualized by separation on a 1.2% agarose gel (Sigma). An approximately 2.5 kb product resulted from amplification of the *Hpal* Vectorette™ library when either ER159 or ER162 were used in combination with the Vectorette™-specific primer, ER70. This fragment represented a 1.9 kb region immediately upstream of the *L. intracellularis* gene encoding AmiB. The PCR product was purified following agarose gel electrophoresis using JETsorb™ kit (GENOMED, Inc., Research Triangle Park, NC) and cloned into pCR2.1 Topo (Invitrogen, Carlsbad, CA) to generate plasmid pER393. The insert was partially sequenced using vector specific primers. The sequence obtained was analyzed by the BLASTx algorithm (Altschul et al., 1990, *J. Mol. Biol.* 215:403-10) and shown to partially encode a protein with similarity to an approximately 85 kDa protein in the GenBank database. The reported proteins from *Neisseria* meningitidis, *Haemophilus influenzae*, and *Pasteurella multocida* were Omp85, protective surface antigen D15, and Oma87, respectively.

[0177] Based on the newly identified sequence of this partial gene fragment (encoding about the C-terminal 2/3 of the Omp100 protein) specific primers ER174 (SEQ ID NO: 46) and ER175 (SEQ ID NO: 47) were designed to obtain additional 5' flanking sequences by a second round of screening the Vectorette™ libraries by PCR amplification. Oligonucleotides ER174 (SEQ ID NO: 46) and ER175 (SEQ ID NO: 47) were used in combination with primer ER70 (SEQ ID NO: 33\ in 50 μl reactions containing 1x PCR Buffer II (Perkin Elmer), 1.5 mM MgCl₂, 200 μM each deoxy-NTP, 50 pMol each primer, and 2.5 U AmpliTaq Gold (Perkin Elmer) thermostable polymerase. Multiple single reactions were performed with 2 μl of the Vectorette™ *Eco*Rl and *Dral* libraries as DNA template. Amplification was carried out as follows: denaturation (95° 9 mm); 35 cycles of denaturation (95° 30 sec), annealing (62° 30 sec), and polymerization (72° 2.5 min); followed by a final extension at 72° for 7 minutes.

[0178] Screening of the *Eco*RI and *Dra*I Vectorette™ libraries by PCR (employing either ER174 or ER175 in combination with ER70) resulted in successful amplification of an approximately 1.4 kb fragment from the *Eco*RI Vectorette™ library. The PCR product was purified following agarose gel electrophoresis using JETsorb™ kit and cloned into pCR2.1 Topo to generate plasmid pER394. Sequence analysis of the insert termini within pER394 using ER175 and a vector specific primer confirmed this fragment was contiguous (e.g. overlapped) with the fragment insert within pER393 and allowed determination of the 5' end of the *omp100* gene. This analysis also indicated the presence of an additional partial ORF having homology to the ATP-binding cassette (ABC) superfamily of transporter proteins. Accordingly, the encoded partial protein was designated ABC1.

[0179] Based on the newly identified nucleotide sequence of this partial gene fragment (encoding about the C-terminal 1/2 of the ABC1 protein) specific primer ER188 (SEQ ID NO: 55) was designed to obtain additional 5' flanking sequence by a third round of screening the Vectorette[™] libraries by PCR amplification. Oligonucleotide ER188 (SEQ ID NO: 55) was used in combination with primer ER70 (SEQ ID NO: 33) in 50 μl reactions containing 1x PCR Buffer II, 1.5 mM MgCl₂, 200 μM each deoxy-NTP, 50 pMol each primer, and 2.5 U AmpliTaq[™] Gold thermostable polymerase. Multiple single reactions were performed with 4 μl of the Vectorette[™] HindIII, DraI, and HpaI libraries as DNA template. Amplification was carried out as follows: denaturation (95° 9 min); 30 cycles of denaturation (95° 30 sec), annealing (60° 30 sec), and polymerization (72° 2.5 min); followed by a final extension at 72° for 7 minutes.

[0180] Screening of the HindIII, Dral, and Hpal Vectorette™ libraries by PCR (employing ER188 in combination with ER70) resulted in successful amplification of an approximately 0.8 kb fragment from the HindIII Vectorette™ library. The PCR product was purified following agarose gel electrophoresis using JETsorb™ kit and cloned into pCR2.1 Topo to generate plasmid pER395. Sequence analysis of the insert termini within pER395 using ER188 and vector specific primers confirmed this fragment was contiguous (e.g. overlapped) with the fragment insert within pER394 and allowed determination of the 5' end of the abc1 gene. An additional partial ORF was identified upstream of the abc1 gene. The encoded protein was designated YcfW based on its homology with the conserved protein, YcfW, found in numerous bacteria.

[0181] The region encoding the remaining portion of the *ycfW* ORF was obtained by screening a Lambda ZAP Express™ phage library created by partial *Tsp509I* digestion of *L. intracellularis* genomic DNA. The phage library was plated onto XL1 -Blue MRF *E. coli* cells in the presence of 10 mM MgSO₄, IPTG, and X-Gal. Clear plaques were picked and phage inserts were amplified using the Expand Long Template PCR System™ as recommended by the supplier (Boehringer Mannheim, Indianapolis, IN). The T3 and T7 phage specific primers were used in PCR conditions consisting of denaturation (94° 2 min); 25 cycles of denaturation (94° 10 sec), annealing (50° 30 sec), and polymerization (68° 6 min); followed by a final extension at 68° for 7 min. Resulting PCR products were en d-sequenced using the T3 and T7 primers and compared to genes in the GeneBank database by BLASTx analysis. One phage, designated clone A21,

contained an approximately 6.1 kb insert encompassing 2.8 kb which overlapped the previously identified *ycfW*, *ABC1*, and *amp100* DNA sequence. Accordingly this clone was used to determine the DNA sequence encoding the N-terminus of the YcfW protein. An additional ORF was identified upstream of the *ycfW* gene. This gene encoded a protein which shares homology with several lysyl-tRNA synthetases and was designated *lysS*.

[0182] The preliminary nucleotide sequence for the *omp100* and C-terminal portion of the *abc1* genes was obtained by sequencing the inserts within pER393 and pER394. Preliminary nucleotide sequence encoding the C-terminal 141 amino acid portion of YcfW and amino-terminal portion of ABC1 was obtained by sequencing the insert within pER395. Preliminary nucleotide sequence encoding the *lysS* and N-terminal portion of the *ycfW* gene was obtained by sequencing the PCR product representing the insert contained in phage clone A21. The primers employed for preliminary sequencing included the vector-specific M13 forward, M13 reverse, phage T3 and T7 primers in addition to ER159 (SEQ ID NO: 37), ER169 (SEQ ID NO: 41), ER170 (SEQ ID NO: 42), ER176 (SEQ ID NO: 48), and ER177 (SEQ ID NO: 49) for pER393; ER175 (SEQ ID NO: 47), ER185 (SEQ ID NO: 52), ER186 (SEQ ID NO: 53), and ER187 (SEQ ID NO: 54) for pER394; ER188 (SEQ ID NO: 55) for pER395; and ER246 (SEQ ID NO: 97), ER254 (SEQ ID NO: 98), ER255 (SEQ ID NO: 99), ER256 (SEQ ID NO: 100), and ER257 (SEQ ID NO: 101) for phage clone A21.

Specific PCR amplification of subgenomic DNA fragments encompassing *L. intracellularis* Region A

[0183] Results of the cloning and preliminary sequencing from the genomic fragments contained in plasmids pER393, pER394, pER395 and phage clone A21 were used to design oligonucleotide primers for the specific amplification of overlapping subgenomic fragments directly from L. intracellularis-infected pig mucosal DNA extracts. DNA extraction was carried out according to the methods described above. This approach was preferred based on the desire to eliminate introduction of sequencing artifacts due to possible mutations arising during the cloning of gene fragments in E. coli. Oligonucleotides ER246 (SEQ ID NO: 97) and ER254 (SEQ ID NO: 98), which flank the lysS and N-terminal 3/4 of ycfW; oligonucleotides ER229 (SEQ ID NO: 73) and ER206 (SEQ ID NO: 66), which flank the abc1 gene; and 25 ER231 (SEQ ID NO: 75) and ER232 (SEQ ID NO: 76), which flank the omp100 gene, were used to specifically amplify this region from the mucosal DNA extract. The lysS gene was amplified in a PCR reaction containing 2 μ l mucosal DNA extract as template, 1x PC2 buffer (Ab Peptides, Inc.), 200 μM each dNTP, 50 pMol each primer, 7.5 U Klen Taq 1 and 0.15 U cloned Pfu thermostable polymerases in a 50 μl final sample volume. Conditions for amplification consisted of denaturation at 94° for 5 minutes followed by 30 cycles of denaturation (94° 1 minute), annealing (55° 30 seconds), and polymerization (72° 3 minuntes). A final extension at 72° for 7 minutes completed the amplification of the targeted 2.6 kb region. The abc1 gene was amplified in triplicate PCR reactions containing 1 μl mucosal DNA extract as template, 1x PC2 buffer, 200 μM each dNTP, 50 pMol each primer, 7.5 U Klen Taq1 and 0.15 U cloned Pfu thermostable polymerases in a 50 µl final sample volume. Conditions for amplification consisted of denaturation at 95° for 5 min followed by 33 cycles of denaturation (94° 1 min), annealing (58° 30 sec), and polymerization (72° 80 sec). A final extension at 72° for 10 minutes completed the amplification of the targeted gene region. The omp100 gene was amplified in quadruplicate PCR reactions containing 2 μl mucosal DNA extract as template, 1x PC2 buffer, 200 μM each dNTP, 50 pMol each primer, 7.5 U Klen Taq1 and 0.15 U cloned Pfu thermostable polymerases in a 50 μl final sample volume. Conditions for amplification consisted of denaturation at 94° for 5 min followed by 35 cycles of denaturation (94° 30 sec), annealing (60° 30 sec), and polymerization (72° 3 min). A final extension at 72° for 7 minutes completed the amplification of the targeted gene region. Following amplification, each of the samples were pooled if appropriate and the specific product was purified by agarose gel electrophoresis prior to direct sequence analysis using DyeDeoxy™ termination reactions on an ABI automated DNA sequencer (Lark Technologies, Inc., Houston, TX).

[0184] Synthetic oligonucleotide primers were used to sequence both DNA strands of the amplified products from L. intracellularis. The primers used for sequence analysis included: AP58.1 (SEQ ID NO: 26), AP58.2 (SEQ ID NO: 27), AP59.1 (SEQ ID NO: 28), AP59.2 (SEQ ID NO: 29), AP59.3 (SEQ ID NO: 30), AP59.4 (SEQ ID NO: 31), AP59.5 (SEQ ID NO: 32), ER159 (SEQ ID NO: 37), ER169 (SEQ ID NO: 41), ER170 (SEQ ID NO: 42), ER175 (SEQ ID NO: 47), ER176 (SEQ ID NO: 48), ER177 (SEQ ID NO: 49), ER185 (SEQ ID NO: 52), ER186 (SEQ ID NO: 53), ER187 (SEQ ID NO: 54), ER188 (SEQ ID NO: 55), ER205 (SEQ ID NO: 65), ER206 (SEQ ID NO: 66), ER217 (SEQ ID NO: 71), ER229 (SEQ ID NO: 73), ER230 (SEQ ID NO: 74), RA138 (SEQ ID NO: 79), RA139 (SEQ ID NO: 80), RA140 (SEQ ID NO: 81), AP182.1 (SEQ ID NO: 83), AP182.2 (SEQ ID NO: 84), AP182.3 (SEQ ID NO: 85), AP182.4 (SEQ ID NO: 86), AP182.5 (SEQ ID NO: 87), AP182.6 (SEQ ID NO: 88), AP182.7 (SEQ ID NO: 89), AP182.8 (SEQ ID NO: 94), AP182.13 (SEQ ID NO: 91), AP182.14 (SEQ ID NO: 92), AP182.11 (SEQ ID NO: 93), AP182.12 (SEQ ID NO: 94), AP182.13 (SEQ ID NO: 95), AP182.14 (SEQ ID NO: 96), ER246 (SEQ ID NO: 97), ER254 (SEQ ID NO: 98), ER255 (SEQ ID NO: 99), ER256 (SEQ ID NO: 100), and ER257 (SEQ ID NO: 101).

The nucleotide sequence of the *L. intracellularis* genomic Region A is listed in SEQ ID NO: 1. The deduced amino acid sequences of the encoded LysS, YcfW, ABC1, and Omp100 proteins within this region are presented in SEQ ID NO: 102, SEQ ID NO: 3, SEQ ID NO: 4, and SEQ ID NO: 5, respectively.

Molecular analysis of the L. intracellularis genes and gene products specified by Region A

[0186] The *L. intracellularis* chromosomal Region A identified upstream of the *amiB* gene encodes proteins designated LysS, YcfW, ABC1, and Omp100 (Fig. 1). These genes are encoded by the same DNA strand and are very closely arranged. This organization suggests these genes may be part of an operon and are likely translationally coupled in the case of LysS/YcfW and ABC1/Omp100. The *IysS* ORF likely initiates from an atypical TTG initiation codon and would extend from nucleotide 165-1745 of SEQ ID NO: 1. This gene encodes a deduced 526 amino acid protein, designated LysS (SEQ ID NO: 102), having a theoretical molecular weight of about 60,628 daltons. The *ycfW* ORF extends from nucleotide 1745-3028 of the reported sequence (SEQ ID NO: 1). This gene encodes a deduced 427 amino acid protein, designated YcfW (SEQ ID NO: 3), having a theoretical molecular weight of about 46,957 daltons. The *abc1* ORF extends from nucleotide 3031 -3738 of SEQ ID NO: 1, and encodes a deduced 235 amino acid protein, ABC1 (SEQ ID NO: 4), having a theoretical molecular weight of about 25,618 daltons. Further downstream but overlapping this ORF by 44 nucleotides is an additional large open reading frame. This ORF, which was designated *omp100*, extends from nucleotide 3695-6388 of SEQ ID NO: 1. The *omp100* gene encodes a deduced 896 amino acid protein which was designated Omp100 (SEQ ID NO: 5). The Omp100 protein has a theoretical molecular weight of about 101,178 daltons.

Similarity of L. intracellularis LysS protein to lysyl-tRNA synthetases

The deduced amino acid sequence of the LysS protein (SEQ ID NO: 102) was compared to other proteins reported in GeneBank by the BLASTp algorithm (Altschul, S.F et al., 1997, Nucleic Acids Res. 25:3389-3402) and aligned by the CLUSTAL W algorithm (Thompson, J.D. et al., 1994, Nucleic Acids Res. 22:4673-4680). As shown in Figure 9, this analysis indicated that LysS shares similarity with members of the cytoplasmic lysyl-tRNA synthetase family. The L. intracellularis LysS protein shares 47% identity with the lysyl-tRNA synthetase protein (Accn. AB012100) from Bacillus stearothermophilus. Consistent with its cytoplasmic location no secretion signal sequence was identified near the predicted N-terminus of this protein.

Similarity of L. intracellularis YcfW and ABC1 proteins to other hypothetical proteins

The YcfW protein shares limited homology with a family of conserved hypothetical proteins approximately [0188] 40-45 kDa in size. Members of this family are reported to be transmembrane or integral membrane proteins. A structural prediction comparison of representative proteins from this family was carried out using TMPred (EMBnet - European Molecular Biology Network; http://www.ch.embnet.org/index.html). The TMpred program makes a prediction of membrane-spanning regions and their orientation. The algorithm is based on the statistical analysis of TMbase, a database of naturally occurring transmembrane proteins. (Hofmann & Stoffel, 1993, Biol. Chem. Hoppe-Seyler 347:166). This analysis indicates that homologs within this protein family have three strong transmembrane domains clustered near the C-terminus of the protein. We have noted an extremely well conserved domain at the very carboxyl-terminal four amino acids (LRYE) of representatives from this family. The observation that the C-te rminal region contains multiple transmembrane domains while the extreme C-terminus is highly conserved suggests a universal functional requirement associated with this region of the YcfW family of homologous proteins. The L. intracellu laris YcfW protein presented in SEQ ID NO: 3 also contains three C-terminal transmembrane domains in addition to the extreme C-terminal amino acids (LRYE). In addition to the three carboxyl domains above, TMPred analysis indic ates that residues 19-44 of the YcfW protein are likely to form a transmembrane region. The amino terminus of YcfW was also examined by the PSORT (Ver. 6.4) computer algorithm using networks trained on known secretion signa. I sequences. This analysis Indicates that residues 29-45 are likely to form a transmembrane region (P. Klein et al., 1985, Biochim. Biophys. Acta, 815:468) which is predicted to act as an uncleavable signal sequence (D. J. McGeoch, Virus Research, 3:271, 1985 and G. von Heijne, Nucl. Acids Res., 14:4683, 1986). As shown in Fig. 2, the 427 am ino acid L. intracellularis YcfW protein shares 32% identity with a 415 residue hypothetical protein (Accn. AJ235272) from Rickettsia prowazekii. The deduced amino acid sequence of the ABC1 protein (SEQ ID NO: 4) was compared to other known pro-

teins reported in GenBank by the BLASTp algorithm. An especially well conserved region (GASGSGKS) was identified near the amino terminus of ABC1. This region corresponds to the nucleotide-binding motif A (P-loop) present in ABC-type transporters. The ABC-type proteins consist of a very large superfamily of proteins which have a wide variety of cellular functions. The majority of these proteins are classified as ABC-type proteins based on regional homology within the nucleotide-binding motif and are generally thought to be involved in cellular transport functions. Figure 3 shows an alignment of ABC1 with that of YcfV from E. coli, (Accn. AE000212) which shares about 45% identical amino acid residues. The E. coli YcfV protein is a probable ABC-type transport protein.

Similarity of L. intracellularis Omp100 protein to 85 kDa proteins

[0190] Examination of the amino terminus of Omp100 indicates that amino acids 1-25 are hydrophobic and positively charged which is characteristic of signal sequences (von Heijm, 1985, *J. Mol. Biol.* 184:99-105). The SignalP (Ver. 1.1) computer algorithm (Nielsen, H., et al., 1997, *Prot. Engineering*, 10:1-6; http://www.cbs.dtv.dk/services/signalP/), using networks trained on known signal sequences, predicted the most likely cleavage site between amino acids 25 and 26. Thus amino acids 1-25 are predicted to be removed from Omp100 during the outer membrane localization process. The Omp100 C-terminal amino acid is predicted to be a phenylalanine residue, a feature consistent with the correct localization of outer membrane proteins (Struyve, M., 1991, *J. Mcl. Biol.* 218:141-148).

[0191] The deduced amino acid sequence of the Omp100 protein (SEQ ID NO: 5) was compared to other known proteins reported in GenBank by the BLASTp algorithm (Altschul, S.F et al., 1997, Nucleic Acids Res. 25:3389-3402) and aligned by the CLUSTAL W algorithm (Thompson, J.D. et al., 1994, Nucleic Acids Res. 22:4673-4680). As shown in Figure 4, this analysis indicated Omp100 shares limited similarity with an approximately 85 kDa protein in the GenBank database (designated U70214). Alignment of the C-terminal ends of Omp100 and this hypothetical protein (YaeT, Accn. U70214 or AE000127) from E. coli indicate these proteins share about 23% identical residues. Other reported proteins include those identified from Shigella flexneri (Oma90), Neisseria meningitidis (Omp85), Haemophilus influenzae (D15), and Pasteurella multocida (Oma87), among others. The NH₂ terminal portion including amino acids 1-139 does not align with any known protein. An additional search of GenBank with the BLASTp algorithm using only the region encompassing amino acids 1-200 of the encoded Omp100 protein failed to detect any known Omp85-like proteins. This data indicates that the amino terminal portion of Omp100 is entirely unique to L. intracellularis.

Example 2: Molecular cloning of L. intracellularis chromosomal Region B

Molecular cloning of genomic Region B encoding PonA, HtrA, HypC, and ORF1 proteins.

[0192] Identification of genomic Region B (shown in Figure 1) was accomplished by a genomic walking process similar to that described for identification of genomic Region A. Screening of the *HindIII*, *EcoRI*, *DraI*, and *HpaI* VectoretteTM libraries was carried out to obtain DNA fragments located adjacent to gene *flgE* from *L. intracellularis* which encodes a protein with homology to the flagellar hook protein of other bacteria. Oligonucleotide primers ER142 (SEQ ID NO: 34), ER153 (SEQ ID NO: 35), and ER158 (SEQ ID NO: 36) were designed based on the known nucleotide sequence 3' of *flgE*. All three primers were designed to bind the (+) strand within this region to allow amplification of DNA located downstream of the gene encoding FlgE.

[0193] For polymerase chain amplification of a fragment of the *ponA* gene, oligonucleotides ER142 (SEQ ID NO: 34), ER153 (SEQ ID NO: 35), and ER158 (SEQ ID NO: 36) were used in combination with a Vectorette™ specific primer (ER70) (SEQ ID NO: 33) in 50 μl reactions containing 1x PCR Buffer II, 1.5 mM MgCl₂, 200 μM each deoxy-NTP, 50 pMol each primer, and 2.5 U AmpliTaq™ Gold thermostable polymerase. Multiple single reactions were performed with 4 μl of the Vectorette™ libraries as DNA template. Amplification annealing temperatures, extension times, and number of cycles varied between experiments and were carried out over the following ranges: denaturation (95° 9 min); 35-40 cycles of denaturation (95° 30 sec), annealing (50-60° 30 sec), and polymerization (72° 2.5-3 min); followed by a final extension at 72° for 7 minutes.

[0194] The amplified products were visualized by separation on a 1.2% agarose get. An approximately 1.2 kb product resulted from amplification of the *Dral* VectoretteTM library when ER158 (SEQ ID NO: 36) was used in combination with the VectoretteTM-specific primer, ER70. Conditions leading to specific amplification of this product included denaturation (95° 9 min); 40 cycles of denaturation (95° 30 sec), annealing (60° 30 sec), and polymerization (72° 2.5 min); followed by a final extension at 72° for 7 minutes. This fragment represented a 1.4 kb region immediately downstream of the *L. intracellularis* gene encoding FIgE. The PCR product was purified following agarose get electrophoresis using a JETsorbTM kit and cloned into pCR2.1 Topo to generate plasmid pER390. The insert was partially sequenced using ER70 and ER158 primers. The sequence obtained was analyzed by the BLASTx algorithm (Altschul, S. F. et al., 1990) and shown to encode a polypeptide with similarity to the amino terminal one half of penicillin-binding proteins in the GenBank database.

[0195] Based on the newly identified sequence of this partial gene, primer ER163 (SEQ ID NO: 40) was designed to obtain additional 3' flanking sequences by a second round of screening the Vectorette™ libraries. Oligonucleotide ER163 (SEQ ID NO: 40) was used in combination with primer ER70 (SEQ ID NO: 33) in 50 µl reactions containing 1x PCR Buffer II, 1.5 mM MgCl₂, 200 µM each deoxy-NTP, 50 pMol each primer, and 2.5 U AmpliTaq Gold thermostable polymerase. Multiple single reactions were performed with 2 µl of the Vectorette™ HindIII, EcoRI and Hpal libraries as DNA template. Amplification was carried out as follows: denaturation (95° 9 min); 30 cycles of denaturation (95° 30 sec), annealing (62° 30 sec), and polymerization (72° 1.5 min); followed by a final extension at 72° for 7 minutes.

[0196] A 2.7 kb fragment was amplified from the *Hin*dIII Vectorette™ library. The PCR product was purified following

agarose gel electrophoresis using JETsorb™ kit and cloned into pCR2.1 Topo to generate plasmid pER392. Sequence analysis of the cloned insert termini using vector specific primers confirmed this fragment was contiguous with the fragment insert within pER390. This analysis also indicated the presence of an additional partial ORF corresponding to approximately the NH₂-terminal 400 residues of the HtrA protein family of serine proteases. Accordingly, the encoded partial protein was designated HtrA.

[0197] A third round of genomic walking was carried out to identify additional sequence within the *htrA* ORF. Specific primer ER173 (SEQ ID NO: 45) was designed based on the known sequence near the 3' end of the insert within pER392. Oligonucleotide ER173 (SEQ ID NO: 45) was used in combination with primer ER70 (SEQ ID NO: 33) in 50 μl reactions as above. Multiple single reactions were performed with 2 μl of the Vectorette[™] *Dral* and *Hpal* libraries-as DNA template. Amplification (denaturation (95° 9 min); 35 cycles of denaturation (95° 30 sec), annealing (62° 30 sec), and polymerization (72° 2.5 min); followed by a final extension at 72° for 7 minutes) resulted in production of a 0.3 kb fragment from the *Dral* library. The PCR product was purified following agarose gel electrophoresis using a JETsorb[™] kit, cloned into pCR2.1 Topo, and the insert sequenced on both strands using vector specific primers. Sequence and BLASTx analysis indicated that the *htrA* ORF remained open through the 3' end of the cloned fragment and that an additional 10 amino acids would be expected to remain before the end of the encoded HtrA protein.

[0198] A final round of genomic walking was carried out to identify the remainder of the *htrA* ORF and 3' flanking region. Specific primer ER189 (SEQ ID NO: 56) was designed based on the known sequence near the 3' end of the *htrA* ORF. Oligonucleotide ER189 (SEQ ID NO: 56) was used in combination with primer ER70 (SEQ ID NO: 33) in 50 μl reactions as above. Multiple single reactions were performed with 4 μl of the Vectorette[™] *Hindlil*, *EcoRI*,and *HpaI* libraries as DNA template. Amplification was carried out as follows: denaturation (95° 9 min); 30 cycles of denaturation (95° 30 sec), annealing (60° 30 sec), and polymerization (72° 2.5 min); followed by a final extension at 72° for 7 minutes. Amplification resulted in production of an approximately 1 kb fragment from the *EcoRI* library. The PCR product was purified following agarose gel electrophoresis using a JETsorb[™] kit and cloned into pCR2.1 Topo to generate pER396. Sequence analysis of the insert termini within pER396 using vector specific primers allowed determination of the 3' end of the *htrA* gene. An additional small ORF was identified downstream of the *htrA* gene. The encoded protein was designated HypC based on its homology with the HypC protein found in other bacteria. Further downstream from *hypC* is an additional partial ORF, designated *orf1*, which is encoded by the opposite DNA strand. This truncated 177 amino acid polypeptide was designated ORF1.

[0199] The preliminary nucleotide sequence for the ponA, htrA, hypC, and C-terminal portion of the orf1 genes was obtained by sequencing the inserts within pER390, pER392 and pER396. The primers employed for preliminary sequencing included the vector-specific M13 forward and M13 reverse primers in addition to ER193 (SEQ ID NO: 59) and ER194 (SEQ ID NO: 60) for pER390; ER171 (SEQ ID NO: 43), ER172 (SEQ ID NO: 44), ER178 (SEQ ID NO: 50), ER179 (SEQ ID NO: 51), ER190 (SEQ ID NO: 57), and ER191 (SEQ ID NO: 58) for pER392; and ER195 (SEQ ID NO: 61) and ER196 (SEQ ID NO: 62) for pER396.

Specific PCR amplification of subgenomic DNA fragments encompassing L. intracellularis Region B

[0200] Results of the cloning and preliminary sequencing from the genomic fragments contained in plasmids pER390, pER392, and pER396 were used to design oligonucleotide primers for the specific amplification of overlapping subgenomic fragments directly from *L. intracellularis*-infected pig mucosal DNA extracts (methods described above for DNA extraction were employed). This approach was preferred based on the desire to eliminate introduction of sequencing artifacts due to possible mutations arising during the cloning of gene fragments in *E. coli*. Oligonucleotides ER228 (SEQ ID NO: 72) and ER190 (SEQ ID NO: 57), which flank the *ponA* gene; oligonucleotides ER207 (SEQ ID NO: 67) and RA134 (SEQ ID NO: 78), which flank the *htrA* gene; and oligonucleotides ER189 (SEQ ID NO: 56) and ER196 (SEQ ID NO: 62), which flank the *hypC* gene were used to specifically amplify this region from the mucosal DNA extract. The endpoints of these fragments overlap thereby allowing specific amplification of subgenomic DNA fragments which are contiguous followed by subsequent confirmation by comparing the terminal nucleotide sequences present in each unique, overlapping DNA fragment. Accordingly, the final sequence represents the complete *L. intracellularis* genomic Region B.

[0201] The overlapping *ponA*, *htrA*, and *hypC* gene regions were amplified in triplicate PCR reactions each containing 1 μl mucosal DNA extract as template, 1x PC2 buffer (Ab Peptides, Inc.), 200 μM each dNTP, 50 pMol each primer, 7.5 U Klen Taq 1 and 0.15 U cloned *Pfu* thermostable polymerases in a 50 μl final sample volume. Conditions for amplification of *ponA* consisted of denaturation at 95° for 5 min followed by 33 cycles of denaturation (95° 30 sec), annealing (62° 30 sec), and polymerization (72° 3 min). Conditions for amplification of *htrA* consisted of denaturation at 94° for 5 min followed by 33 cycles of denaturation (95° 30 sec), annealing (58° 30 sec), and polymerization (72° 3 min). Conditions for amplification of *hypC* consisted of denaturation at 94° for 5 min followed by 33 cycles of denaturation (95° 30 sec), annealing (62° 30 sec), and polymerization (72° 80 sec). A final extension at 72° for 7 minutes completed the amplification of each of the targeted gene regions. Following amplification, each of the samples were pooled

separately and the specific product was purified by agarose gel electrophoresis prior to direct sequence analysis using DyeDeoxy™ termination reactions on an ABI automated DNA sequencer (Lark Technologies, Inc., Houston, TX).

[0202] Synthetic oligonucleotide primers were used to sequence both DNA strands of the amplified products from L. intracellularis. The primers used for sequence analysis included: AP55.1 (SEQ ID NO: 10), AP55.2 (SEQ ID NO: 11), AP55.3 (SEQ ID NO: 12), AP55.4 (SEQ ID NO: 13), AP55.5 (SEQ ID NO: 14), AP55.6 (SEQ ID NO: 15), AP55.7 (SEQ ID NO: 16), AP55.8 (SEQ ID NO: 17), AP55.9 (SEQ ID NO: 18), AP55.10 (SEQ ID NO: 19), AP55.11 (SEQ ID NO: 20), AP56.1 (SEQ ID NO: 21), AP56.2 (SEQ ID NO: 22), AP56.3 (SEQ ID NO: 23), AP57.1 (SEQ ID NO: 24), AP57.2 (SEQ ID NO: 25), ER158 (SEQ ID NO: 36), ER163 (SEQ ID NO: 40), ER171 (SEQ ID NO: 43), ER172 (SEQ ID NO: 44), ER173 (SEQ ID NO: 45), ER178 (SEQ ID NO: 50), ER179 (SEQ ID NO: 51), ER189 (SEQ ID NO: 56), ER190 (SEQ ID NO: 57), ER191 (SEQ ID NO: 58), ER193 (SEQ ID NO: 59), ER194 (SEQ ID NO: 60), ER195 (SEQ ID NO: 61), ER196 (SEQ ID NO: 62), ER203 (SEQ ID NO: 63), ER204 (SEQ ID NO: 64). ER207 (SEQ ID NO: 67), ER208 (SEQ ID NO: 68), ER213 (SEQ ID NO: 69), ER228 (SEQ ID NO: 72), RA133 (SEQ ID NO: 77), RA134 (SEQ ID NO: 78), and RA171 (SEQ ID NO: 82).

[0203] The nucleotide sequence of the *L. intracellularis* genomic Region B is listed in SEQ ID NO: 2. The deduced amino acid sequences of the encoded PonA, HtrA, HypC, and ORF1 proteins within this region are presented in SEQ ID NO: 6, SEQ ID NO: 7, and SEQ ID NO: 8, and SEQ ID NO: 9, respectively.

Molecular analysis of the L. intracellularis genes and gene products specified by Region B

[0204] The L. intracellularis chromosomal Region B identified downstream of the flgE gene encodes proteins designated PonA, HtrA, HypC, and a partial "ORF1" protein (Fig. 1). A portion of the figE ORF is presented here and extends from nucleotide 1-125 (SEQ ID NO: 2). The ponA ORF extends from nucleotide 252-2690 of SEQ ID NO: 2, and encodes a deduced 812 amino acid protein, PonA (SEQ ID NO: 6), having a theoretical molecular weight of about 90,263 daltons. An alternative in-frame translation initiation codon is present at nucleotide 276 which, if utilized, would encode a slightly smaller 804 amino acid protein having a theoretical molecular weight of about 89,313 daltons. The htrA ORF extends from nucleotide 2981-4315 of SEQ ID NO: 2, and encodes a deduced 474 amino acid protein, HtrA (SEQ ID NO: 7), having a theoretical molecular weight of about 50,478 daltons. The small hypC ORF extends from nucleotide 4581-4829 of SEQ ID NO: 2, and encodes a deduced 82 amino acid protein, HypC (SEQ ID NO: 8), having a theoretical molecular weight of about 8,888 daltons. Further downstream and transcribed in the opposite orientation is an additional open reading frame. This ORF, which was designated "orf1", extends from nucleotide 4912-5445 at the 3' end of SEQ ID NO: 2. This ORF remains open through the 3' end of SEQ ID NO: 2 and thus encodes the C-terminal 177 amino acids of a truncated protein having a theoretical molecular weight of at least about 20,345 daltons. As shown in Fig. 8, the encoded ORF1 protein (SEQ ID NO: 9) shares limited homology with a 205 amino acid hypothetical protein encoded by gene "MJ1123" (Accn. U67555) from the Methanococcus jannaschii genome. 35

Similarity of L. intracellularis HypC protein to hydrogenase maturation proteins

The HypC protein shares homology with the hyp/hup family of hydrogenase maturation proteins. Hydrogenase, which catalyzes the reversible oxidation of molecular hydrogen, is involved in many relevant anaerobic processes where hydrogen is oxidized or produced (Adams, M.W.W., et al., 1980, *Biochem. Biophys. Acta* 594:105-176). The HypC protein is required for the maturation of catalytically active hydrogenase isozymes in *E. coli*. The precise role of HypC in this process is unknown but hydrogenase maturation involves nickel insertion, protein folding, C-terminal proteolytic processing, membrane integration, and reductive activation (Lutz, S., et al., 1991, *Mol. Microbiol.* 5:123-135; Przybyla, A.E., et al., 1992, *FEMS Microbiol. Rev.* 88:109-136). The HypC protein is 41% identical to the *Desultovibrio gigas* 82 amino acid HynD protein (Accn. AJ223669, as shown in Figure 7) and 39% identical to the 75 amino acid HypC protein from *Rizobium leguminosarum*.

Similarity of L. Intracellularis PonA protein to penicillin binding proteins

The ponA ORF encodes a deduced 812 amino acid protein, having a theoretical molecular weight of about 90,263 daltons. An alternative in-frame methionine codon is present which encodes a slightly smaller 804 amino acid protein having a theoretical molecular weight of about 89,313 daltons. Similar in-frame methionine codons have been identified in other characterized ponA ORF's. For example, PonA homologs from Neisseria flavescens (Accn. AF087677), N. gonorrhoeae (Accn. U72876), and N. meningitidis (Accn. U80933) contain amino-terminal in-frame methionine codons separated by 8, 6, and 6 codons, respectively. As with L. intracellularis, the neisserial ponA genes are preceded by undiscernable ribosome binding sites thus further complicating prediction of the true initiating methionine. N-terminal sequencing of the N. gonorrhoeae FA19 PonA protein indicated the second methionine was the preferred start site in this strain (Ropp et al., 1997, J. Bacteriol. 179:2783-2787). The upstream methionine codon was

used as the putative initiation site for the encoded PonA protein from L. intracellularis.

[0207] A structural prediction of the PonA protein was carried out using TMPred. The TMpred program makes a prediction of membrane-spanning regions and their orientation (K. Hofmann & W. Stoffel, 1993. TMbase - A database of membrane spanning proteins segments. *Biol. Chem.* Hoppe-Seyler 347:166). This analysis indicates that PonA has a strong transmembrane domain at the NH₂-terminus of the protein. The amino terminus of PonA was examined by the PSORT (Ver. 6.4) computer algorithm using networks trained on known signal sequences. This analysis indicates that residues 16-32 are likely to form a transmembrane region (P. Klein et al., 1985, *Biochim. Biophys. Acta*, 815:468) which is predicted to act as an uncleavable signal sequence (D. J. McGeoch, *Virus Research*, 3:271, 1985 and G. von Heijne, *Nucl. Acids Res.*, 14:4683, 1986). Thus the amino terminus of PonA is predicted to anchor the protein to the bacterial inner membrane, which is similar to the method of localization of other penicillin-binding proteins.

The PonA protein shares homology with Class A high-molecular-mass penicillin-binding proteins (PBP's) identified in other bacteria. Penicillin-binding proteins are bacterial cytoplasmic membrane proteins involved in the final steps of peptidoglycan synthesis. The Class A proteins generally exhibit two types of enzymatic activities: the glycosyltransferase, which polymerizes the glycan strand and the transpeptidase, which cross-links these strands by their peptide side chains. These reactions are catalyzed either on the outer surface of the cytoplasmic membrane or further outside and the major fraction of the proteins involved in peptidoglycan synthesis is therefore localized in the periplasm. The deduced amino acid sequence of the PonA protein (SEQ ID NO: 6) was compared to other known proteins reported in GenBank by the BLASTp algorithm (Altschul, S.F et al., 1997, supra) and aligned by the CLUSTAL W algorithm (Thompson et al., 1994, supra). As shown in Figure 5, this analysis indicated PonA is most similar to a penicillinbinding protein from Neisseria flavescens (Accn. AF087677). PonA shares features characteristic of class A highmolecular-mass PBPs. The sequence including amino acids 124-134 (RQGGSTITQQV) corresponds to a highly conserved consensus amino acid sequence known as the QGAST box (Popham et al., 1994, J. Bacteriol. 176:7197-7205) found in all class A high-molecular-mass PBPs. Within the C-terminal half of PonA, three regions can be found that are highly conserved in all members of the penicilloyl serine transferase superfamily. These regions include the SXXK tetrad containing the active site serine at residues 507-510 (SAFK), the SXN triad at residues 565-567 (SRN), and the KT(S)G tetrad at residues 688-691 (KTG). These motifs are thought to be brought close together in the folded protein to form the transpeptidase domain active-site pocket that interacts with β -lactam antibiotics.

Similarity of L. intracellularis HtrA protein to periplasmic serine protease proteins

[0209] Examination of the amino terminus of HtrA indicates that amino acids 1-26 are hydrophobic and positively charged which is characteristic of signal sequences (von Heijm, 1985, J. Mol. Biol. 184:99-105). The PSORT computer algorithm (Nakai, K., 1991, PROTEINS: Structure, Function, and Genetics 11: 95-110), using networks trained on known signal sequences indicates that residues 1-26 likely function as a typical signal sequence and predicts the most likely cleavage site between amino acids 26 and 27. Thus amino acids 1-26 are predicted to be removed from HtrA during the maturation process.

[0210] The deduced amino acid sequence of the HtrA protein (SEQ ID NO: 7) was compared to other known proteins reported in GenBank by the BLASTp algorithm (Altschul, S.F et al., 1997, above) and aligned by the CLUSTAL W algorithm (Thompson, J.D. et al., 1994, supra). This analysis indicated HtrA belongs to the large HtrA/DegP family of periplasmic serine proteases. The reported proteins include those identified from E. coli, Salmonella typhimurium, Camplylbacter jejuni, Haemophilus influenzae, Brucella melitensis, Brucella abortus, Chlamydia trachomatis, Yersinia enterocolitica, Borrelia burgdorferi, and Bacillus subtilis, among others. In some instances the HtrA homolog is referred to as a heat shock protein and has been shown by deletion analysis to be required for bacterial survival at elevated temperatures or for survival of intracellular pathogens. In other cases an HtrA homolog is not induced by temperature but is expressed in response to other physiological stress. Several HtrA homologs have been shown to possess serine protease activity and in a number of cases is important for bacterial virulence and/or intracellular survival, for example resistance to high temperature, hydrogen peroxide, oxidative and osmotic stress.

[0211] Alignment of the *L. intracellularis* HtrA protein with its most similar relative from *Pseudomonas aeruginosa* (Accn. #U32853) indicates the two proteins share 40% identical amino acid residues (as shown in Figure 6). Based on alignment of the *L. intracellularis* HtrA protein with other serine proteases, especially well conserved residues including Histidine-109, Aspartic acid-143, and the active-site Serine-217 are predicted to form the catalytic triad of residues which are highly conserved in bacterial and mammalian serine proteases. A number of HtrA homologs contain a carboxy-terminal RGD motif while others have been shown to contain an RGN motif. The *L. intracellularis* HtrA protein contains a similar motif at residues 458-460 (RNG). The RGD motif has been identified as a cell attachment site for mammalian adhesion proteins (Ruoslahti, E. et al., 1986, *Cell* 44:517-518). The HtrA/DegP family of serine proteases are induced during a range of stress responses and during infection by *L. intracellularis*, surface expression of HtrA may occur as part of a stress response mechanism. Other intracellular heat shock proteins have been shown to become surface expressed under physiological stress conditions and have been implicated as adh esion factors (Ensgraber, M. et

al,, 1992, Infect. Immun. 60:3072-3078 and Hartmann, E. et al., 1997, Infect. Immun. 65:1729-1733).

Analysis of the htrA promoter region and induction in response to temperature

The gene arrangement for L. intracellularis Region A and Region B differ with regard to the extent of intergenic spacing between the encoded proteins. Unlike Region A the ORF's within Region B are more distantly separated. For example, the flgE, ponA, htrA, and hypC genes are separated by approximately 125, 200, and 265 nucleotides between the respective open reading frames. The 200 bp region immediately upstream of htrA was examined in more detail to find a promoter region, particularly since several HtrA protein homologs have been shown to be induced in response to a number of different environmental signals including temperature, oxidative, and osmotic stress. Examination of the nucleotide sequence of SEQ ID NO: 2 upstream of htrA indicated a promoter located about nucleotide 2797-2802 (TTGATA; -35 region) and nucleotide 2824-2829 (TATAAT; -10 region). These two hexamers are separated by a 21 nucleotide space and share near perfect homology to consensus sigma 70 type promoters. Other promoter elements may exist in this region which control htrA expression in response to various environmental signals. Plasmid pER434, which contains the htrA ORF and htrA promoter region imparts a temperature-dependent phenotype to E. coli host cells grown at either 30°C or 37°C. Thus, the region upstream of htrA can be recognized as a likely functional promoter in response to temperature. It should therefore be possible to use the htrA promoter to operably control expression of heterologous proteins in E. coli and other organisms in response to temperature. The presence of other promoter elements that control expression in response to other environmental signals would allow those other signals to be used to control expression.

Example 3: Preparation of plasmids and deposit materials

Plasmids containing DNA fragments encompassing L. intracellularis Region A

[0213] Plasmids were prepared containing the *L. intracellularis* genomic region representing the *lysS*, *ycfW*, *abc1*, and *omp100* genes. A 2.6 kb fragment encompassing the *lysS* gene and a portion of the *ycfW* gene was amplified using primers ER246 (SEQ ID NO: 97) and ER254 (SEQ ID NO: 98) while a 0.87 kb fragment encompassing a portion of the *ycfW* gene and complete *abc1* gene fragment was amplified using primers ER229 (SEQ ID NO: 73) and ER206 (SEQ ID NO: 66). These fragments were amplified as described in Example 1 under "Specific PCR amplification of subgenomic DNA fragments encompassing *L. intracellularis* Region A*. The 2.6 kb and 0.87 kb DNA fragments were isolated by extraction with spin chromotography (QIAquick™) and inserted into the TA cloning site of pCR2.1 Topo. Single sequence extension reactions utilizing vector-specific sequencing primers confirmed the endpoints of the cloned fragments, and revealed that the genes encoding LysS and YcfW in plasmid pT068 and YcfW and ABC1 in plasmid pER438 were in the opposite orientation relative to the lactose promoter.

[0214] A 2.97 kb DNA fragment containing the *omp100* gene was amplified by PCR employing specific 5' and 3' primers ER187 (SEQ ID NO: 54) and ER170 (SEQ ID NO: 42). PCR reactions were carried out in triplicate and contained 1 μl DNA extract as template, 1x PCR Buffer II, 1.5 mM MgCl₂, 200 μM each deoxy-NTP, 50 pMol each primer, and 2.5 U AmpliTaq Gold thermostable polymerase in a 50 μl final sample volume. Conditions for amplification consisted of denaturation at 95° for 9 min followed by 33 cycles of denaturation (95° 30 sec), annealing (62° 30 sec), and polymerization (72° 3 min). A final extension at 72° for 7 minutes completed the amplification of the target gene region. Following amplification, each of the triplicate samples were pooled and the specific product was isolated by extraction with spin chromotography (QIAquick™) and inserted into the TA cloning site of pCR2.1 Topo in the opposite orientation relative to the lactose promoter. This plasmid construct was designated pER440.

Plasmids pER438 and pER440 were introduced into *E. coli* TOP 10 cells (Invitrogen, Carlsbad, CA). The resulting strains, designated Pz438 and Pz440, were deposited with the ATCC (10801 University Blvd, Manassas, VA, pT068 was introduced into *E. coli* TOP10 cells and the resulting strain was deposited with the ATCC on July 14, 2000 and assigned accession number PTA-2232.

Plasmids containing DNA fragments encompassing L. intracellularis Region B

[0216] Plasmids were prepared containing the *L. intracellularis* genomic region representing the *ponA*, *htrA*, and *hypC* genes. The ponA, *htrA*, and *hypC* gene fragments were amplified as described above in Example 2, in the section entitled "Specific PCR amplification of subgenomic DNA fragments encompassing *L. intracellularis* Region B" using primers ER228 (SEQ ID NO: 72) and ER190 (SEQ ID NO: 57), which flank the *ponA* gene; primers ER207 (SEQ ID NO: 67) and RA134 (SEQ ID NO: 78), which flank the *htrA* gene; and primers ER189 (SEQ ID NO: 56) and ER196 (SEQ ID NO: 62), which flank the *hypC* gene. The resulting 2.98 kb fragment containing *ponA* was purified following

agarose gel electrophoresis using a JETsorb™ kit and cloned into pCR2.1 Topo to generate plasmid pER432. The resulting 1.72 kb fragment containing *htrA* was isolated by extraction with spin chromotography (QIAquick™) and inserted into the TA cloning site of pCR2.1 Topo to generate plasmid pER434. The resulting 0.98 kb fragment containing *hypC* and additional flanking nucleotides encoding the C-terminal 94 amino acids of ORF1 was isolated by extraction with spin chromotography (QIAquick™) and inserted into the TA cloning site of pCR2.1 Topo to generate plasmid pER436. Single sequence extension reactions utilizing vector-specific sequencing primers confirmed the endpoints of the cloned fragments, and revealed that the genes encoding PonA and HypC were in the opposite orientation relative to the lactose promoter. The HtrA gene was cloned in the same orientation relative to the lactose promoter and cells containing such plasmids exhibited an unstable phenotype at 37°C which was relieved when growth was maintained at 30°C.

[0217] Plasmids pER432, pER434 and pER436 were introduced into *E. coli* TOP10 cells (Invitrogen, Carlsbad, CA). The resulting strains, designated Pz432, Pz434, and Pz436 were deposited with the ATCC (10801 University Blvd, Manassas, VA, 20110, USA) on September 9, 1999 and assigned the accession numbers PTA-635, PTA-636, and PTA-637, respectively.

Example 4: Expression of recombinant HtrA and Omp100 proteins in E. coli

Plasmid expression vectors

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[0218] The expression vector used for production of recombinant HtrA and Omp100 was pET-28b (+) (Novagen, Inc., Madison, WI). The coding sequences for the HtrA and Omp100 proteins were amplified from *L. intracellularis*-infected pig mucosal DNA extract. The PCR products were purified following agarose gel electrophoresis using a JET-sorb™ kit and cloned into pCR2.1 Topo to generate plasmids pRL001 (HtrA) and pER415 (Omp100). Specific PCR primers used to amplify the HtrA ORF included ER208 (SEQ ID NO: 68) and RA133 (SEQ ID NO: 77). Primer ER208 was designed to introduce an *Ndel* site (CATATG) while deleting the HtrA signal sequence. The HtrA insert present in pRL001 was subcloned into pET-28b (+) following digestion with *Ndel* and *EcoRI*. The resulting expression plasmid, designated pER405, was sequenced at both 5' and 3' ends of the inserted fragment and confirmed to encode an inframe fusion with the vector encoded 6x His leader. Therefore the predicted amino terminal sequence of the encoded protein consisted of the sequence MGSSHHHHHHHSSGLVPRGSHM encoded by the vector followed immediately by the sequence ALPNFVP beginning at Alanine-27 of the HtrA open reading frame.

[0219] Specific PCR primers used to amplify the Omp100 ORF included ER216 (SEQ ID NO: 70) and RA138 (SEQ ID NO: 79). Primer ER216 was designed to introduce an *Ncol* site (CCATGG) while deleting the Omp100 signal sequence. In addition, ER216 specified a leader peptide, termed a "protective peptide" which protects recombinant proteins from proteolytic degradation, based on information from Sung et al., 1986, Proc. Natll. Acad. Sci. USA 83:561-565; Sung et al., 1987, Meth. Enzymol. 153:385-389; and U.S. Patent 5,460,954, which references are incorporated herein by reference. The protective peptide consisting of the amino acid sequence MG TTTTTSL was encoded by the 5' proximal nucleotide sequence of ER216. The Omp100 insert present in pER415 was subcloned into pET-28b (+) following digestion with *Ncol* and *EcoRl*. The resulting expression plasmid, designated pRL029, was sequenced at both 5' and 3' ends of the inserted fragment and confirmed to encode an in-frame fusion with the protective peptide leader. Therefore the predicted amino terminal sequence of the encoded protein consisted of the sequence MGTTTTTTSL specified by the 5' proximal nucleotide sequence of ER216 followed immediately by the sequence ASKDDPSIV beginning at Alanine-26 of the Omp100 open reading frame.

Expression of recombinant proteins

[0220] The pET-28b (+) based expression vectors pER405 and pRL029, encoding recombinant HtrA and Omp100, respectively, were introduced into the expression host *E. coli* BL21(DE3). This expression host has the genotype F, ompT hsdS_B (r_B·m_B·) gal dcm (DE3) (Novagen, Inc.) which allows high level transcription of cloned genes driven by the IPTG-inducible phage T7 promoter. The *E. coli* transformants were propagated in SB#2 medium (2.4% yeast extract, 1.2% tryptone, 0.5% K₂HPO₄, 0.25% KH₂PO₄, .014% MgSO₄) containing 50 µg/mI kan amycin sulfate in a 5 L BioFlow 3000 fermentor (New Brunswick Scientific, Edison, NJ) at 30-37 °C until A₆₂₅ was 2.5-30.1. Recombinant protein expression was obtained following induction with 1 mM IPTG for 1-4.5 h.

[0221] Wet cells of *E. coli* expressing recombinant HtrA were lysed by homogenization at 10,000 psi (2 passes) followed by centrifugation. The pellet, which contained HtrA, was washed with 2x RIPA/TET which was in a 5:4 ratio. 2x RIPA is 20 mM Tris (pH 7.4), 0.3 M NaCl, 2.0% sodium deoxycholate, and 2% (v/v) Igep at CA-630[™] (Sigma). TET is 0.1 M Tris (pH 8.0), 50 mM EDTA, and 2% (v/v) Triton X-100. The washed pellet was then solubilized in 8 M Urea, 10 mM Tris, 0.1 M NaH₂PO₄, pH 7.0. The solubilized protein was diluted 2 fold in 8 M Urea, 1 O mM Tris, 0.1 M NaH₂PO₄, pH 7.0 and applied onto a Ni NTA column (QIAGEN, Santa Clarita, CA). The desired prote in was eluted off this column by

reduction of pH into 8 M Urea, 10 mM Tris, 0.1 M NaH_2PO_4 , pH 4.5. The final pooled fractions were dialyzed against 4 M Guanidine HCl, 50 mM Tris, pH 6.5 and then step dialyzed to 2 M Guanidine HCl, 25 mM Tris, pH 6.5. The final product was filtered by 0.22 μ M filtration. The protein concentration was 0.56 mg/ml with an estimated visual purity of 70% by SDS-PAGE.

[0222] Wet cells of *E. coli* expressing recombinant Omp100 were lysed with lysozyme and sonication in the presence of Benzonase™ (Benzonase™ (EM Industries Inc, Hawthorne, New York)), to facilitate DNA degradation, followed by centrifugation. The pellet, which contained Omp100, was washed twice with 2 M Urea, 50 mM Tris, 10 mM EDTA, ugation. The pellet was resuspended with 6 M Urea, 50 mM Tris (pH 8.0) followed by centrifugation. The pellet was washed with 2x RIPA/TET which was in a 5:4 ratio and the washed pellet was then solubilized in 8 M Urea, 50 mM Tris (pH 8.0). 25 mM DTT was added to the solubilized protein and further diluted 2:1 with 8 M Urea, 25 mM DTT, 50 mM Tris (pH 8.0). The diluted solubilized protein was applied onto a Q-Sepharose column equilibrated with 8 M Urea, 25 mM DTT, 50 mM Tris (pH 8.0). Recombinant Omp100 was eluted in a linear gradient of 0-1 M NaCl in 8 M Urea, 25 mM DTT, 50 mM Tris (pH 8.0). The pooled fractions were dialyzed against 6 M Guanidine HCl, 10 mM DTT, 50 mM Tris (pH 8.0) and then step dialyzed to 4 M Guanidine HCl, 6.7 mM DTT, 33.3 mM Tris (pH 8.0). The final product was filtered by 0.22 μM filtration and frozen at 70°C. The purified Omp100 protein was then thawed and centrifuged (16,000 rpm, 60 min) and the supernatant was subjected to 0.22 μM filtration again to remove insoluble particles and aggregates. The protein concentration was 1.08 mg/ml with an estimated visual purity of 80% by SDS-PAGE.

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					85					90					95	
5	Ile	Ser	Leu	Arg 100		Phe	Gly	Lys	Val 105	Ile	Phe	Phe	His	11e	Met	Asp
	Arg	Ser	Gly 115	Arg	Ile	Gln	Cys	Туг 120	Ala	Ser	Arg	Glu	Asn 125	мet	G1 y	Glu
10	Glu	Ala 130	Phe	Ser	Thr	Phe	Lys 135	Lys	Phe	Asp	Ile	Gly 140	Asp	Ile	Val	Gly
15	Val 145		Gly	Lys	Leu	Phe 150	Arg	Thr	Lys	Met	Gly 155	Glu	Leu	The	Leu	Asn 160
	Cys	Ser	Thi	Ile	Thr 165	Leu	Leu	Ala	Lys	Ser 170	Phe	Arg	Ser	Leu	Pro 175	G1 u
20	Lys	Hls	Asn	Gly 180	Leu	Thr	Asn	Ile	Glu 185	Leu	Arg	туг	Arg	Gln 190	Arg	Tyr
25	Ile	Asp	Leu 195	Ile	Val	Asn	Pro	Lys 200	Thr	Arg	Asp	Ile	Phe 205	Arg	Lys	Arg
	Ser	Lys 210	Ile	lle	Hıs	Glu	11e 215	Arg	Ala	Phe	Leu	Glu 220	Glu	Asn	GJ À	Phe
30	Ile 225	Glu	Val	Glu	Thr	Pro 230	Ile	Leu	Gln	Pro	Ile 235	Pro	Gly	Gly	Ala	Met 240
35	Ala	Arg	Pro	Phe	Thr 245	Thr	His	Asn	Asn	Ala 250	Met	Asp	Met	Thr	Leu 255	Туr
	Met	Arg	Ile	Ala 260	Pro	Glu	Leu	Tyr	Leu 265	Lys	Arg	Leu	Leu	Val 270		Gly
40	₽h€	Glu	Lys 275	Leu	Phe	Glu	Leu	Asn 280	Arg	ser	Phe	Arg	Asn 285	Glu	Gly	lle
	Ser	11e 290	Gln	His	Asn	Pro	Glu 295	Phe	Thr	Met	Cys	Glu 300	Phe	Туr	Trp	Ala
45	Туг 305	Ala	Thir	Tyr	Leu	Asp 310	Leu	Met	Glu	Leu	Thr 315	Glu	Glu	Met	Phe	Ala 320
50	Tyr	Leu	Thr	Lys	Lys 325	Ile	Cys	G1 y	Thr	Met 330	Thr	lle	Ser	Туг	335	n Gly
	Asn	Thr	Ile	Asp	Phe	Thr	Pro	Gly	Thr	Trp	Gln	Lys	туг	Thi	Phe	e His

				3 4	0				34	5				35	0	
5	G1	u Se	er Le	eu Gl SS	u Ly	s Ile	e Gl	y G1 36	у Ні О	s Se	r Pro	o G1	u Ph		r Ası	n Asn
10	Ph	€ G1 37	u Ly 'O	'S Va	l Sei	r Glu	и Тул 375	Ile	e Ly:	s Gl	u His	38)		Ly:	s Val	Leu
	Th. 38:	r Th	r As	p Ly:	s Ile	Gly 390	Lys	Let	Glr	Ala	1 Lys 395	Let	ı Ph∈	e Asp) Lei	Asp 400
15	Va)	. G1	u Ası	n Lys	Leu 405	Ile	Gln	Pro	Thr	Phe 410	lle	Туг	His	Туг	Pro	
20	Asp	116	e Sej	Pro 420	Leu	Ser	Lys	Lys	Asn 425	Lys	Asp	Asn	Pro	Glu 430		Thr
	Asp	Arg	Phe 435	Glu	Leu	Phe	Ile	Ala 440	Gly	Lys	Glu	Ile	Ala 445	Asn	Ala	Phe
25	Ser	Glu 450	Leu	Asn	Asp	Pro	Ile 455	Asp	Gln	Arg	Leu	Arg 460	Phe	Glu	Glu	Gln
30	Val 465	Leu	G1 u	Lys	Ala	Arg 470	Gly	Asp	Glu	Glu	Ala 475	Cys	Pro	Met	Asp	Glu 480
25	Asp	Туr	Leu	Arg	Ala 485	Leu	Glu	Туг	Gly	Met 490	Pro	Pro	Ala	Ala	Gly 495	Glu .
35	Gly	lle	Gly	11e 500	Asp /	Arg :	Leu '	Val	Met 505	Leu	Leu	Thr	Asp	Ser 510	Pro	Ser
40	lle ,	Arg	Glu 515	Val	lle :	Leu 1		Pro :	Leu	Leu .	Arg '		Glu 525	Arg		

45 Claims

- 1. An isolated polynucleotide molecule comprising a nucleotide sequence that is selected from the group consisting of
 - a) a nucleotide sequence encoding *L. intracellularis* HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein; b) a nucleotide sequence that is a substantial part of said nucleotide sequence encoding said *L. intracellularis* HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein; and
 - c) a nucleotide sequence that is homologous to the nucleotide sequence of a) or b).
- The isolated polynucleotide molecule of claim 1 comprising said nucleotide sequence encoding said L.
 intracellularis HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein.
 - 3. The isolated polynucleotide molecule of claim 1 comprising a nucleotide sequence consisting of the open reading frame selected from the group consisting of the open reading frame of SEQ ID NO: 1 from about nt 165 to about nt

1745, the open reading frame of SEQ ID NO: 1 from about nt 3031 to about nt 3738, the open reading frame of SEQ ID NO: 1 from about nt 3695 to about nt 6385, the open reading frame of SEQ ID NO: 2 from about nt 252 to about nt 2690. the open reading frame of SEQ ID NO: 2 from about nt 2891 to about nt 4315, and the open reading frame of SEQ ID NO: 2 from about nt 4581 to about nt 4829.

- A polynucleotide molecule comprising a nucleotide sequence of greater than 20 nucleotides having promoter activity and found within SEQ ID NO: 2 from about nt 2691 to about nt 2890, or its complement.
- A recombinant vector comprising the polynucleotide molecule of claim 1.
- 6. The recombinant vector of claim 5 consisting of a plasmid selected from the group consisting of plasmid pER432 containing the ponA gene (ATCC accession number PTA-635), plasmid pER434 containing the htrA gene (ATCC accession number PTA-636), plasmid pER436 containing the hypC gene (ATCC accession number PTA-637), plasmid pT068 containing the lysS and ycfW genes (ATCC accession number PTA-2232), plasmid pER438 containing the ycfW and abc1 genes (ATCC accession number PTA-638), and plasmid pER440 containing the Omp100 gene (ATCC accession number PTA-639).
- A transformed host cell comprising the recombinant vector of claim 5.
- A polypeptide produced by the transformed host cell of claim 7.
 - A genetic construct comprising a polynucleotide molecule that can be used to alter a Lawsonia gene, comprising: (a) a polynucleotide molecule comprising a nucleotide sequence that is otherwise the same as a nucleotide sequence of a htrA, ponA, hypC, lysS, ycfW, abc1, or omp100 gene, or that is otherwise the same as a nucleotide sequence that is homologous thereto, or a substantial portion of said nucleotide sequence, but which nucleotide sequence further comprises one or more mutations capable of altering the htrA, ponA, hypC, lysS, ycfW, abc1, or omp100 gene; or (b) a polynucleotide molecule comprising a nucleotide sequence that naturally flanks in situ the ORF of said htrA, ponA, hypC, lysS, ycfW, abc1, or omp100 gene, or a nucleotide sequence that is homologous to said flanking sequence; such that transformation of a Lawsonia cell with the genetic construct of (a) or (b) results in altering the htrA, ponA, hypC, lysS, ycfW, abc1, or omp100 gene.
 - 10. A transformed cell comprising the genetic construct of claim 9.
 - 11. An isolated polypeptide, said polypeptide being selected from the group consisting of:
 - (a) L. intracellularis HtrA, PonA, HypC, YcfW, ABC1, or Omp100 protein;
 - (b) a polypeptide having an amino acid sequence that is homologous to that of said L. intracellularis HtrA, PonA, HypC, YcfW, ABC1, or Omp100 protein;
 - (c) a polypeptide consisting of a substantial portion of said L. intracellularis HtrA, PonA, HypC, YcfW, ABC1, or Omp100 protein or of said polypeptide having an amino acid sequence that is homologous to that of said L. intracellularis HtrA, PonA, HypC, YcfW, ABC1, or Omp100 protein;
 - (d) a fusion protein comprising the protein or polypeptide of (a), (b) or (c) fused to another protein or polypep-
 - (e) an analog or derivative of the protein or polypeptide of (a), (b), (c) or (d).
 - 12. The isolated polypeptide of claim 11 wherein said L. intracellularis protein has an amino acid sequence that comprises an amino acid sequence selected from the group consisting of SEQ ID NO: 3, SEQ ID NO: 4, SEQ ID NO: 5, SEQ ID NO: 6, SEQ ID NO: 7, SEQ ID NO: 8, and SEQ ID NO: 102.
- 13. The isolated polypeptide of claim 11 comprising a polypeptide that consists of said L. intracellularis HtrA, PonA, 50 HypC, LysS, YcfW, ABC1, or Omp100 protein having between 1 and 10 amino acids inserted, deleted, or substituted, including combinations thereof.
- 14. A substantially pure polypeptide comprising an epitope of HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein that is specifically reactive with anti-Lawsonia antibodies. 55
 - 15. An isolated polypeptide comprising an amino acid sequence encoded by the polyn ucleotide molecule of claim 1.

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- An isolated antibody that specifically reacts with L. intracellularis HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 protein.
- 17. A live attenuated vaccine comprising the transformed cell of claim 10.
- 18. A killed cell vaccine comprising transformed cells of claim 10 in killed form.
- 19. An immunogenic composition comprising an immunogenically effective amount of the polypeptide of claim 11 in combination with a pharmaceutically acceptable carrier.
- 20. An immunogenic composition comprising an immunogenically effective amount of a polynucleotide molecule comprising a nucleotide sequence encoding the polypeptide of claim 11 in combination with a pharmaceutically acceptable carrier.

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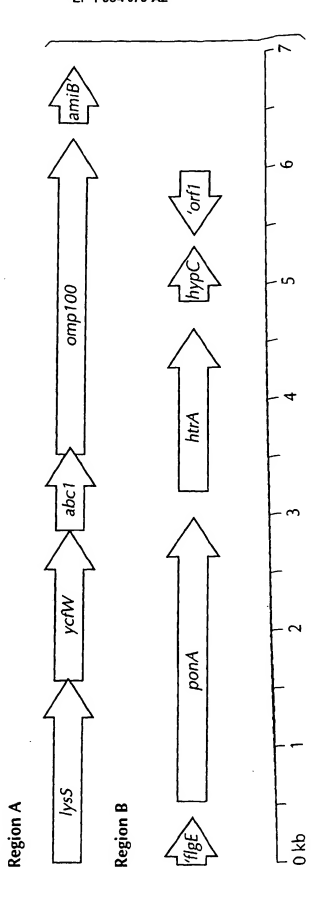
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Li_Ycfw AJ23527	2 MINNNFSFNIAFRYFRAKKNEKFVSIIAAFSLYCVMICVAALIVERKINGSTIDIRDKI	57 60
Li_Ycfw AJ235272		.58
Li_Ycfw AJ235272	MISSPHGVKGLILRGIDPSSAQNVISMLSHLTKGNLEDLIPKVLGTPDGIIIGNELAQRL 1	77 58
Li_Ycfw AJ235272	NVTIGSRVNLLSPTGQKTSSGFQPRIRPLIVTGIFHTGMFEYDTSLAFTSLNAARELLGL 2 GVTVGEKLRLISPNSVSTAFGSIPRSKEFOLLA IENGGREVAN GENERALDEL 2	37 18
Li_Ycfw AJ235272	PHYTISGIEVSIHDVYOANYITWOLOOFI CHNESUPCURCURCURCURCURCURCURCURCURCURCURCURCURC	97 77
Li_Ycfw AJ235272	AMVVLIGSFSIVTTLIMLVMEKTRDIAILTSMGATSQMIRRIFILQGTIIGIVGTLLGYL 35 SLIITVAAFNIISNLFMLVKDKTSDIAILRTMGASTKQIMVIFIYNGMFIGLLGTTLGVI 33 :::::::::::::::::::::::::::::::::::	57 57
AJ235272	LGITLALLLQKYQFIKLPPGVYTIDHLPVLLNWLDIFIIGTSAMLLCFFATL 40 LGVTFSYNIQTIKNYLERITGIKIFEAAIYFLYSLPSKVKTDDIILITSLSIILCFLATI 39 **:*:: :* :: :::**::*:::::*:::::*:::::::	-
Li_Ycfw AJ235272	YPAHQAARLQPIEGLRYE 427 YPSYRASKLNPVDALRYE 415 **:::*::*::****	

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Li_OMP10 U70214	0 MTKRLNIFLLLLCNILYCNIIANAASKDDPSIVVLPFQINGSSNDEELQTELPMLLATA	60
Li_OMP10 U70214	0 LKNKGFRVIPNKSALNLLYKQNISQLNISTAKKVAQQLHADYVVYGSFNQTGENFSIDSR	120
Li_OMF10 U70214	0 LIDSTGVASARPLYIEKPKFNELNIAVTELAERISNGLIKKNTIADVRIHGLKVLDPDVIMAMKKLLIASLLFSSATVYG-AEGFVVKDIHFEGLQRVAVGAA :::* ** ::. * : :: *:::.**: :	180 42
070214	<pre>DETRITINKGDHTDHAKINAEIKKIWELGYFSDVSASIEESGEGRLLVFTVQEKPKITDVV LLSMPVRTGDTVNDEDISNTIRALFATGNFEDVRVLRDGDTLLVQVKERPTIASIT * : : : * * : : * * . * : : * * . * . *</pre>	240 98
Li_OMP10(U70214	VQGSKAVSIDNILAAMSSKKGSVISDRLLSQDIQK-ITDLYRKEGYYLAEVNYEIKEK FSGNKSVKDDMLKQNLEASGVRVGESLDRTTIADIEKGLEDFYYSVGKYSASVKAVVTPL	297 158
Li_OMP100 U70214	P-RNRVDLKLVFQEGVSAEIQQINIVGNHAGTTDELISHFQLRDEVPWWNVVGDRKYQ	355 215
Li_OMP100 U70214	EEYLERDSIAISAYAMNHGYVDIQVASPEVTFNEKGIVITFRVKEGKRYKIGKIDFKG KQKLAGDLETLRSYYLDRGYARFNIDSTQVSLTPDKKGIYVTVNITEGDQYKLSGVEVSG :: * * ::: * ::: * ::: * ::: * :: * ::	413 275
Li_OMP100 U70214	DLIETNEQLLKVTKIDDHKNYEQYFSLSVMQDDVKALTDFYSDYGYAFAEVDLETTKNEE NLAGHSAEIEQLTKIEPGE-LYNGTKVTKMEDDIKKLLGRYGYAYPRYQSMPEINDA :* :::***: : :::***: * ***************	473 331
Li_OMP100 U70214	DATIOVTFLIDKKQKVFLRRIIVEGNTRTRDNVILRELRLADGDLFNGQHLRRSNECLNR DKTVKLRVNVDAGNRGYVRKIRFEGNDTSKDAVLRREMRQMEGAWLGSDLVDQGKERLNR	533 391
Li_OMP100 U70214	LGYFNQVDTDTLPT-GKDDEVDLLVKVQEARTGAITGGVGYSTHSKFGVSGSISERNLWG LGFFETVDTDTQRVPGSPDQVDVVYKVKERNTGSFNFGIGYGTESGVSFQAGVQQDNWLG **: *: *****	592 451
Li_OMP100 U70214	KGYILSIEGFISSKSSSLDLSFTNPRVYDTDFGFSNNIYTLRDEWDDFRKKTYGD TGYAVGINGTKNDYQTYAELSVTNPYFTVDGVSLGGRLFYNDFQADDADLSDYTNKSYGT .**: *: *: *: *: *: *: *: *: *: *: *: *:	647 511
Li_OMP100 U70214	TIRLFHPIGEYSSIFVGYRIDQYRLYDI-PSTAPRSYLDYQGKNISSVVSGGFT DVTLGFPINEYNSLRAGLGYVHNSLSNMQPQVAMWRYLYSMGEHPSTSDQDNSFKTDDFT : * .**.**.*:	700 571
070214	FDSTDSRERPSKGHIAKLIVEYGGGGLGGNDN-FFKPIAELQGFYSISRSKNHIIHW FNYGWTYNKLDRGYFPTDGSRVNLTGKVTIPGSDNEYYKVTLDTATYVPIDDDHKWVVLG *: :: : * : : * : * : *	756 631
070214	RTRAGAAYKNSKKPVPVFDRFFIGGIDSIRGYDTEDLAPKDP RTRWGYGDGLGGKEMPFYENFYAGGSSTVRGFQSNTIGPKAVYFPHQASNYDPDYDYECA	691
L1_OMP100 U70214	TQDGAKDLCKSDDAVGGNAMAVASLEFITPTPFISDKYANSVRTSFFWDMGTVWDTNWDS	844 751
0,0214	SQYSGYPDYSDPSNIRMSAGIALQWMSPLGPLVFSYAQPFKKYDGDKAEQFQFNIGKT ::* *:* *:* *:* * * *:* * * *:* * *:* * *:* * *:* * *:* * *	895 809
Li_OMP100 I U70214 V	F 896 W 810	

Li_PonA AF087677	-MKQVISFDMKKFFLNIVIFCFGIILLSIIGLIGLYFWVSRDLPNITKLNDYRPALVTTV MIKKIITTCMGLNNGLALFGVGLIAIAILVTYPKLPSLDSLQHYKPKLPLTI :*::*: *	59 52
Li_PonA AF087677	LARDGTLIGYIYREKRFLIPLSEMSPFLPKAFLAAEDAEFYEHEGVNPLAIIRAFLINLQ YSSDGQVIGVYGEQRREFTKIDDFPKILKDAVIAAEDKRFYDHWGVDVWGVARAVIGNVM : ** : ** : : : : : : : : : : * : * : *	119 112
Li_PonA AF087677	SGTTRQGGSTITQQVIKRLLLSPERSYERKIKEAILAYRLEKYLSKDEILTIYLNQTFLG AGGVQSGASTITQQVAKNFYLSSERSFTRKFNEALLAYKIEQSLSKDKILELYFNQIYLG :**.****** *.: **.***: **::**:**: ***::** :**:	179 172
Li_PonA AF087677	AHSYGVEAAARTYFAKHAKDLSLAECALLAGLPQAPSRYNPYKDPEAAKIRQRYALRRLH QRAYGFASAAQTYFNKNVNDLTLAEAAMLAGLPKAPSAYNPIVNPERAKLRQAYILNNML ::**: *** * * : : * * : * * : * * * : * * * * : * * * * : * * * * : * * * * : * * * * : * * * * : * * * * : * * * * : * * * * : * * * * : * * * * : * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * * : * * * * * * : * * * * * * : * * * * * : * * * * * : * * * * * * * : * * * * * : * * * * * * : * * * * * : * * * * * : * * * * * * : * * * * * * : * * * * * * : * * * * * : * * * * * : * * * * * : * * * * * * : * * * * * : * * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * * : * * * * * : * * * * * * : * * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * : * * * * : * * * * : * * * * : * * * * : * * * * * : * * * * * : * * * * : * * * * : * * * * * : * * * * : * * * * : * * * * * : * * * * : * * * * : * * * * : * * * * : * * * * : * * * * : * * * * * : * * * * * : * * * * * : * * * * : * * * * : * * * * * : * * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * : * * * * * * : * * * * * : * * * * * * : *	239 232
Li_PonA AF087677	DVGWITQAEYEEALQEPLYFSSMKEGLGAESSWYMEEVRKQLVSFLSKENISQYGIVLPL EEGMITLQQRDQALKEELHYERFVQNIDQSALYVAEMARQELFEK : * * * : ::**:* *:::::::::: * .*::* *	299 277
Li_PonA AF087677	YGEDALYELGFTIQTAMDPQAQLVAYDVLRNGLENFSKRQGWKGP1EHISSTMIQHY YGEDA-YTQGFKVYTTVDTAHQRVATEALRKVLRNFDRGSSYRGAENYIDLSKSDNVEET	356 336
Li_PonA AF087677	LENATFTPEKLDGGAWAKAIVSKVSQEGAEVFLSSIYKGFVSVETMGWARKPNPEVRSAY VSQYLSTLYTVDKMIPAVVLEASRKGVQIQLPSGRKVTLNNHALGFAARAVN	416 388
Li_PonA AF087677	CAPIKDARSVLNPGDIIWVSGVGPDSTHRYSSKTLDTSKPIPLALQOBPQIQGALISIEP NEKMGDDRIRRGSVIRVKGSG-DTFTVVQEPDTQGALVSLDA : * : * : * : * : * : * : * : * : * : *	476 429
Li_PonA AF087677	NTGDVIAMIGGYEFGKSQFNRAVQAMRQPGSAFKPIVYSAALDHDYTSATMVLDAPIVEF KTGAVRALVGGYDYHSKTFNRATQAMRQPGSTFKPFIYSAALAKGMTASTMINDAPISLP :** * *::**:: ****.*********** : . *::**: ****	536 489
Li_PonA AF087677	MESGDIWRPGNYEKNFKGPMLFSNALALSRNLCTVRIAQSIGLPAVIERAKALGFNGN GKGANGKAWNPKNSDGRYAGYITLRQALTASKNMVSIRILMSIGIGYAQQYIQRFGFKPS : .*. *.* * : : : **: *: : : ** ***: . : : : **:	594 549
Li_PonA AF087677	-FPEFFSISLGAVEVTPIRLVNAYTAFANGGNLATPRFILSIKDSNN-TVIYRQEIEQHP EIPASLSMALGAGETTPLRIAEGYSVFANGGYKVSAHVIDKIYDSQGRLRAQMQPLVAGE :* : : : : * * * * * * * * * * * * * *	652 609
Li_PonA AF087677	VISPQNAYIMASLLKNVVNIGTARKAKVLERP-LAGKTGTTNGEHDAWFIGFTPYL NAPQAIDPRNAYIMYKIMQDVVRVGTARGAATLGRSDIAGKTGTTNDNKDAWFVGFNPNV .*.*:**** .::::**.:** .: : : ****** : : : ******* : : : ****** : : : : ******	707 669
Li_PonA AF087677	VTGVYVGNDHPQTLGKDGTGAVAALPIFTEYSKVVLKKYPESDFPVPDGITFASIDTQTG VTAVYIGFDKPRSMGRAGYGGTIAVPVWVEYIGFALKGTSVKPMKAPEGVVTNGGEVYMR **.**: *:*::*: * * *:*:** *: .	767 729
Li_PonA AF087677	NRATANSTNSVVLPFYVGTVPEYFDSKDNEVNTIERGED ERMTTSSDLALDNSGIRPRPTQPARRAVPNENRRAESNTAPAREESDETPVLPSNTGNN :* **.	806 789
Li_PonA AF087677	LLKQFF 816 NRQQLDSLF 798	

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Li_Htr/ U32853	A MFCKLKVIICITLMFIITVVPTIAESALPNFVPLVKDASKAVVNISTEKKIPRGRTE -MHTLKRCMAAMVALLALSLAMTARAELPDFTPLVEQASPAVVNISTRQKLPDRAMARGQ : .** :: :: *: **:*.***::** ******.:*: * : *:	57 59
Li_HtrA U32853	FPMEMFRGLPPGFERFFEQFEPKGPDSQIHKQRSLGTGFIISSDGYIVTNNHVIEGA LSIPDLEGLPPMFRDFLERSIPQVPRNPRGQQREAQSLGSGFIISNDGYILTNNHVVADA ::: :.*** *: *: *: *: *: *: *: *: *: *: *: *:	114 119
Li_HtrA U32853	DSVRVNLEGTSGKEESLPAEVIGRDEETDLALLKVKSKDSLPYLIFGNSDTMEVGEWVLA DEILVRLSDRSEHKAKLIGADPRSDVAVLKIEAKN-LPTLKLGDSNKLKVGEWVLA *.: *.*. * * *::** * .:*:*:*: ** * ::*:********	174 174
Li_HtrA U32853	IGNPFGLGHTVTAGILSAKGRDIHAGPFDNFLQTDASINPGNSGGPLINMSGQVVGINTA IGSPFGFDHSVTAGIVSAKGRSLPNESYVPFIQTDVAINPGNSGGPLLNLQGEVVGINSQ **.***:.*:******::******:	234 234
Li_HtrA U32853	IMAS-GQGIGFAIPSSMADRIIEQLKTNKKVSTGWIGVTIQDVDTNTAKALGLSQAKG IFTRSGGFMGLSFAIPIDVALNVADQLKKAGKVSTGWLGVVIQEVNKDLAESFGLDKPSG *:: * *:*** : * * * * * * * * * * * * *	291 294
Li_HtrA U32853	ALVGSVVPGDPADKAGLKVGDIVTQADGKQIDSASSLLKAIATKPPFSVVKLKVWRDGKS ALVAQLVEDGPAAKGGLQVGDVILSLNGQSINESADLPHLVGNMKPGDKINLDVIRNGQR ***:*** *.**:***: .:*:.*:.* :: .:*:.*:	351 354
Li_HtrA U32853	KSI,SMAVCSI,DDDDDRRTA CMCA DOARD COMPA COMPA DO TOTAL	407 402
Li_HtrA U32853	U-DGPAAVTGLR PCDVTTHI DNV AUTCOVI TO DVA VAL DVAD	467 470
Li_HtrA U32853	SLTEDSN 474 KLAE 474 .*:*	

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60 59 MCLAIPAKVIELLDNDIIRATVGDSTTILTVSGMLLPEPVTVGDYIIVHAGFAIHKLEAT MCLAIPAR-IETIENGVATCRVGASDTFVKASLLLLEGQAGPGDYLVVHAGFALRKMDVK ****** 82 82 Li_Hypc AJ223629 Li_HypC AJ223629

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	EFQLGALDDLPFEDESFNYASLVTILEYVEDPKKILAEAFRVASDG- 46 SKEMAKIAEKRGIKVIIAKGEDLPFKDEEFDFFLINTVLEFAENPKKMIEEAKRVLKRGG 120 :.:::::*******************************	-IIVGFTNKWSINHIINSTLQLLHKKPKKDSQWVSPWQLIRLTKQLYPECRIYCRSTLLG 105 KIIIGIIDRDSFLGKMYEEKKQKSK-FYKDANFLSAKEVIEMLKELGFKN 169 **;*; *; *; *; *; *; *; *; *; *; *; *; *	PKRTWDVTSSWSKLNRIILSFPIG TYVGMRIEKRPKPTLTPLLLKAKEQAVNVYNALS 153 IKATQTIFKEIDKVDKVEVKEGYGEGGFVAISAEKI
Li_ORF1	Li_ORF1	Li_ORF1	Li_ORF1
U67555	U67555	U67555	U67555

PEATSTIQHNRTNK

Li_ORF1 U67555

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ii_LysS AB012100_LysS	LIQKKKSHPPIKLATKSPHVSYFKPLLESLAEKNELNEVIKNCVVKSC MSHEELNDQLRVRREKLK
Li_Lyss AB012100_Lyss	EFVKEHYAGMLRAEYEAYSASELESLDEIFACAGRIISLRSFGKVIFFHIMDRSGRIGCY 120 RFERTHKAEELFELYGDLSKEELEEQQIEVAVAGRIMTKRGMGKAGFAHIQDVTGQIQIY 90
Li_LysS AB012100_LysS	ASRENMGEEAFSTFKKFDIGDIVGVNGKLFRTKMGELTLNCSTITLLAKSFRSLPEKHNG 180 VRQDDVGEQQYELFKISDLGDIVGVRGTMFKTKVGELSIKVSSYEFLTKALRPLPEKYHG 150 ::::**:::****************************
Li_LysS AB012100_LysS	LTNIELRYRQRYIDLIVNPKTRDIFRKRSKIIHEIRAFLEENGFIEVETPILQPIPGGAM 240 LKDIEQRYRQRYLDLIMNPESKKTFITRSLIIQSMRRYLDSHGYLEVETPMMHAVAGGAA 210
Li_Lyss AB012100_Lyss	ARPFTTHNNAMDMTLYMRIAPELYLKRLLVGGFEKLFELNRSFRNEGISIQHNPEFTMCE 300 ARPFITHHNALDMTLYMRIAIELHLKRLIVGGLEKVYEIGRVFRNEGISTRHNPEFTMLE 270 **** **: **: ******* **: **: **: **: * *** *** *** *** *
Li_LysS AB012100_LysS	FYWAYATYLDLMELTEEMFAYLTKKICGTMTISYQGNTIDFTPGTWQKYTFHESLEKIGG 360 LYEAYADFRDIMKLTENLIAHIATEVLGTTKIQYGEHLVDLTP-EWRRLHMVDAIKEYVG 329 :* *** : *:*:**::*::::: ** . *
Li_Lyss AB012100_Lyss	HSPEFYNNFEKVSEYIKEHGEKVLTTDKIGKLQAKLFDLDVENKLIQPTFIYHYPTDISP 420 VDFWRQMSDEEARELAKEHGVEVAPHMTFGHIVNEFFEQKVEDKLIQPTFIYGHPVEISP 389
Li_LysS AB012100_LysS	LSKKNKDNPEVTDRFELFIAGKEIANAFSELNDPIDQRLRFEEQVLEKARGDEEACFMDE 480 LAKKNPDDPRFTDRFELFIVGREHANAFTELNDPIDQRQRFEEQLKEREQGNDEAHEMDE 449 *:*** *:* :***************************
Li_Lyss AB012100_Lyss	PLLRTER 526 PQMRHK- 494

(11) EP 1 094 070 A3

(12)

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(51) Int Cl.7: C07K 14/205, C12N 15/31

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- (30) Priority: 22.10.1999 US 160922 P
- (83) Declaration under Rule 28(4) EPC (expert solution)
- (71) Applicant: Pfizer Products Inc. Groton, Connecticut 06340 (US)
- (72) Inventor: Rosey, Everett Lee, Pfizer Central Research Groton, Connecticut 06340 (US)
- (74) Representative: Eddowes, Simon et al Urquhart-Dykes & Lord, 30 Welbeck Street London W1G 8ER (GB)
- (54) Lawsonia intracellularis proteins, and related methods and materials
- (57) Isolated polynucleotide molecules contain a nucleotide sequence that encodes a *L. intracellularis* HtrA, PonA, HypC, LysS, YcfW, ABC1, or Omp100 pro-

tein, a substantial portion of the sequences, or a homologous sequence. Related polypeptides, immunogenic compositions and assays are described.

FIG. 1

EP 1 094 070 A3



EUROPEAN SEARCH REPORT

Application Number EP 00 30 9125

 	DOCUMENTS CONSIDERED TO BE RE			
Category	Citation of document with indication, where appropr of relevant passages			ASSIFICATION OF THE PLICATION (InlCI.7)
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	DALE C JANE H ET AL: "Identifica sequencing of the groE operon and genes of Lawsonia intracellularis phylogeny." MICROBIOLOGY (READING), vol. 144, no. 8, August 1998 (1998) pages 2073-2084, XP002156652 ISSN: 1350-0872	flanking : Use in		
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	ISSN: 0021-9193 abstract; figure 7 *			
	he present search report has been drawn up for all claim			
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LACK OF UNITY OF INVENTION SHEET B

Application Number EP 00 30 9125

The Search Division considers that the present European patent application does not comply with the requirements of unity of invention and relates to several inventions or groups of inventions, namely:

1. Claims: 1-3, 5-20 (all partially), 4 (completely)

A nucleotide sequence encoding L. intracellularis HtrA protein (SEQ ID NO:2, nt 2891 - nt 4315) and subject-matter relating thereto.

2. Claims: 1-3, 5-20 (all partially)

A nucleotide sequence encoding L. intracellularis PonA protein (SEQ ID NO:2, nt 252 - nt 2690) and subject-matter relating thereto.

3. Claims: 1-3, 5-20 (all partially)

A nucleotide sequence encoding L. intracellularis HypC protein (SEQ ID NO:2, nt 4581- nt 4829) and subject-matter relating thereto.

4. Claims: 1-3, 5-10, 12-18 (all partially)

A nucleotide sequence encoding L. intracellularis LysC protein (SEQ ID NO:1, nt 165- nt 1745) and subject-matter relating thereto.

5. Claims: 1-2, 5-20 (all partially)

A nucleotide sequence encoding L. intracellularis YcfW protein (SEQ ID NO:1, nt 1745 - nt 3028) and subject-matter relating thereto.

6. Claims: 1-3, 5-20 (all partially)

A nucleotide sequence encoding L. intracellularis ABC1 protein (SEQ ID NO:1, nt 3031- nt 3738) and subject-matter relating thereto.

7. Claims: 1-3, 5-20 (all partially)

A nucleotide sequence encoding L. intracellularis Omp100 protein (SEQ ID NO:1, nt 3695 - nt 6385) and subject-matter relating thereto.

Article 82 EPC stipulates that the European patent application shall relate to one invention only or to a group so linked as to form a single general inventive concept. Where a group of inventions is claimed in one and the same European patent application, the requirement of unity of invention shall be fulfilled only when there is a technical relationship among those inventions involving one or more of the same or corresponding "special technical features", i.e. technical features that define a novel and inventive contribution over the prior art (Rule 30 EPC).

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LACK OF UNITY OF INVENTION SHEET B

Application Number EP 00 30 9125

The Search Division considers that the present European patent application does not comply with the requirements of unity of invention and relates to several inventions or groups of inventions, namely:

The present application discloses the identification of seven nucleotide sequences encoding seven different Lawsonia intracellularis proteins (HtrA, PonA, HypC, LysC, YcfW, ABC1 and Omp100).

Nucleotide sequences encoding various Lawsonia intracellularis proteins are already known from the prior art (see e.g. Dale C.J.H. et al., Microbiology (1998), 144, 2073-2084 or W09720050).

The problem underlying the present application can, thus, be seen as the provision of further nucleotide sequences encoding alternative Lawsonia intracellularis proteins.

The solutions as disclosed and claimed in the present application can be summarised as the provision of the 7 sequences and the polypeptides they encode (SEQ ID Nos:1-8 and 102).

Due to the fact that nucleotide sequences encoding Lawsonia intracellularis proteins are already known from the prior art, due to the essential differences between the primary structures of the sequences claimed and due to the fact that no other technical feature can be distinguished which in light of the prior art could be regarded as a special, common technical feature, the Search Division is of the opinion that there is no single inventive concept underlying the plurality of different inventions of the present application in the sense of Rule 30 EPC.

Consequently there is lack of unity, and the different inventions not belonging to a common inventive concept are formulated as the different subjects in the communication pursuant to Rule 46 EPC.

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4

ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 00 30 9125

This annex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

17-01-2001

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